

STIC Search Report Biotech-Chem Library

STIC Database Tracking Number: 180215

TO: Nyeemah Grazier

Location: REM-5B29&5C18

Art Unit: 1626 March 10, 2006

Case Serial Number: 10/765267

From: P. Sheppard

Location: Remsen Building

Phone: (571) 272-2529

sheppard@uspto.gov

Search Notes

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V'- mars

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Scientific and Technical Information Center

Prula

SEARCH REQUEST FORM

| Requester's Full Name: Nyeewah (3002 Date: 2206 Art Unit: 1626 Phone Number: 2-878 (Serial Number: 10/765, 267 Location (Bldg/Room#): Lew 5029 (Mailbox #): 5C18 Results Format Preferred (circle): 1 APER DISK *********************************** |
|---|
| To ensure an efficient and quality search, please attach a copy of the cover sheet, claims, and abstract or fill out the following: |
| Title of Invention: Medhods, Mixtures & Kits Restaining to Analyte Determine Inventors (please provide full names): Pappin et al. |
| Earliest Priority Date: 1/27/04 |
| Search Topic: Please provide a detailed statement of the search topic, and describe as specifically as possible the subject matter to be searched. Include the elected species or structures, keywords, synonyms, acronyms, and registry numbers, and combine with the concept or utility of the invention. Define any terms that may have a special meaning. Give examples or relevant citations, authors, etc., if known. |
| *For Sequence Searches Only* Please include all pertinent information (parent, child, divisional, or issued patent numbers) along with the appropriate serial number. |
| Prease Seavel. the compound of Claim 71. |
| $G_1=C_1N_1O$ V |
| W= NH, NR', NRZ, PR', PRZ, O or S J=H, deuterium(D), R', OR', SR', NHR', N(R')z, Fluoring Cl, Br, I |
| Z = O,S, NH, KE'NR' |
| LG = scheded from the group: |
| $\begin{cases} x' - x' \\ y' - x' \end{cases} = \begin{cases} x' - q - x' \\ y' - q - x' \end{cases}$ |

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Grazier 10_765267 - - History

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(FILE 'HOME' ENTERED AT 10:18:35 ON 10 MAR 2006)

FILE 'REGISTRY' ENTERED AT 10:18:44 ON 10 MAR 2006

L1 STR

L5 82 SEA SSS FUL L1

L6

L16

L7 46 SEA SUB=L5 SSS FUL L6

STR

FILE 'HCAPLUS' ENTERED AT 10:24:11 ON 10 MAR 2006

L8 29 SEA ABB=ON PLU=ON L7

D STAT QUE L8

D IBIB ABS HITSTR L8 1-29

FILE 'REGISTRY' ENTERED AT 10:27:29 ON 10 MAR 2006

L9 36 SEA ABB=ON PLU=ON L5 NOT L7

FILE 'HCAPLUS' ENTERED AT 10:27:38 ON 10 MAR 2006

L10 18 SEA ABB=ON PLU=ON L9

L11 17 SEA ABB=ON PLU=ON L10 NOT L8

D STAT QUE

D IBIB ABS HITSTR L11 1-17

L12 103 SEA ABB=ON PLU=ON ("PAPPIN D"/AU OR "PAPPIN D J"/AU OR "PAPPIN D J C"/AU OR "PAPPIN DARRYL"/AU OR "PAPPIN DARRYL J"/AU OR "PAPPIN DARRYL JOHN

CECIL"/AU OR "PAPPIN DARYL"/AU) NOT (L8 OR L11)

9 SEA ABB=ON PLU=ON ("BARTLET JONES M"/AU OR "BARTLET JONES MICHAEL"/AU) NOT (L8 OR L11)

D STAT QUE L13

D IBIB ABS L13 1-9

L14 97 SEA ABB=ON PLU=ON L12 NOT L13

L15 92 SEA ABB=ON PLU=ON L14 AND PD=<JANUARY 28, 2004

38 SEA ABB=ON PLU=ON L15 AND ANALY?

D STAT QUE

D IBIB ABS L16 1-38

FILE 'BEILSTEIN' ENTERED AT 10:35:47 ON 10 MAR 2006

L17 7 SEA SSS FUL L6

L18 6 SEA ABB=ON PLU=ON L17 NOT L7

D STAT QUE

D CN BRN MF FW STR RX 1-6

FILE HOME

FILE REGISTRY

Property values tagged with IC are from the ZIC/VINITI data file provided by InfoChem.

STRUCTURE FILE UPDATES: 8 MAR 2006 HIGHEST RN 876273-86-8 DICTIONARY FILE UPDATES: 8 MAR 2006 HIGHEST RN 876273-86-8

New CAS Information Use Policies, enter HELP USAGETERMS for details.

TSCA INFORMATION NOW CURRENT THROUGH January 6, 2006

Please note that search-term pricing does apply when conducting SmartSELECT searches.

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Grazier 10_765267 - - History

* The CA roles and document type information have been removed from * the IDE default display format and the ED field has been added, * effective March 20, 2005. A new display format, IDERL, is now * available and contains the CA role and document type information. *

Structure search iteration limits have been increased. See HELP SLIMITS for details.

REGISTRY includes numerically searchable data for experimental and predicted properties as well as tags indicating availability of experimental property data in the original document. For information on property searching in REGISTRY, refer to:

http://www.cas.org/ONLINE/UG/regprops.html

FILE HCAPLUS

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FILE COVERS 1907 - 10 Mar 2006 VOL 144 ISS 12 FILE LAST UPDATED: 9 Mar 2006 (20060309/ED)

New CAS Information Use Policies, enter HELP USAGETERMS for details.

This file contains CAS Registry Numbers for easy and accurate substance identification.

FILE BEILSTEIN
FILE LAST UPDATED ON JANUARY 17, 2006

FILE COVERS 1771 TO 2005.
FILE CONTAINS 9,428,406 SUBSTANCES

>>>PLEASE NOTE: Reaction Data and substance data are stored in separate documents and can not be searched together in one query. Reaction data for BEILSTEIN compounds may be displayed immediately with the display codes PRE (preparations) and REA (reactions). A substance answer set retrieved after the search for a chemical name, a compounds with available reaction information by combining with PRE/FA, REA/FA or more generally with RX/FA. The BEILSTEIN Registry Number (BRN) is the link between a BEILSTEIN compound and belonging reactions. For mo detailed reaction searches BRNs can be searched as reaction partner BRNs Reactant BRN (RX.RBRN) or Product BRN (RX.PBRN).<<<

| *** | **** | ******* | ********* | **** | **** | **** | ****** |
|-----|------|-----------|--------------|------|------|------|--------|
| >>> | FOR | SEARCHING | PREPARATIONS | SEE | HELP | PRE | <<< |

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Grazier 10_765267 - - History

* PLEASE NOTE THAT THERE ARE NO FORMATS FREE OF COST. * SET NOTICE FEATURE: THE COST ESTIMATES CALCULATED FOR SET NOTICE * ARE BASED ON THE HIGHEST PRICE CATEGORY. THEREFORE; THESE * ESTIMATES MAY NOT REFLECT THE ACTUAL COSTS. * FOR PRICE INFORMATION SEE HELP COST ******************

NEW

- * PATENT NUMBERS (PN) AND BABS ACCESSION NUMBERS (BABSAN) CAN NOW BE SEARCHED, SELECTED AND TRANSFERRED.
- * NEW DISPLAY FORMATS ALLREF, ALLP AND BABSAN SHOW ALL REFERENCES, ALL PATENT REFERENCES, OR ALL BABS ACCESSION NUMBERS FOR A COMPOUND AT A GLANCE.

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=> fil hcaplus FILE 'HCAPLUS' ENTERED AT 10:24:11 ON 10 MAR 2006 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2006 AMERICAN CHEMICAL SOCIETY (ACS)

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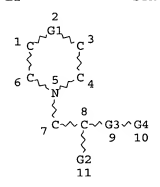
FILE COVERS 1907 - 10 Mar 2006 VOL 144 ISS 12 FILE LAST UPDATED: 9 Mar 2006 (20060309/ED)

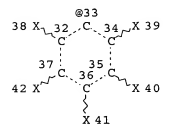
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This file contains CAS Registry Numbers for easy and accurate substance identification.

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VAR G1=C/N/O VAR G2=O/S/N VAR G3=O/S VAR G4=13/19/26/33 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM

DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

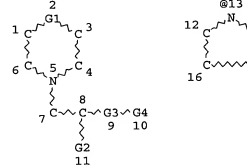
RING(S) ARE ISOLATED OR EMBEDDED

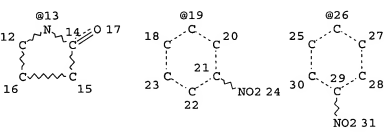
NUMBER OF NODES IS 42

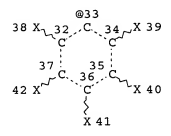
STEREO ATTRIBUTES: NONE

L5 82 SEA FILE=REGISTRY SSS FUL L1

L6 STR







VAR G1=C/N/O
VAR G2=O/S/N
VAR G3=O/S
VAR G4=13/19/26/33
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC 1

NUMBER OF NODES IS 42

STEREO ATTRIBUTES: NONE

L7 46 SEA FILE=REGISTRY SUB=L5 SSS FUL L6 L8 29 SEA FILE=HCAPLUS ABB=ON PLU=ON L7

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=> d ibib abs hitstr 18 1-29

L8 ANSWER 1 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2005:592130 HCAPLUS

DOCUMENT NUMBER: 143:115574

TITLE: Preparation of isotopically enriched N-substituted

piperazines

INVENTOR(S): Pappin, Darryl J. C.; Pillai, Sasi; Coull, James M.

PATENT ASSIGNEE(S): Applera Corp., USA

Grazier 10 765267

U.S. Pat. Appl. Publ., 29 pp. SOURCE:

CODEN: USXXCO

DOCUMENT TYPE: LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PA | PATENT NO. | | | | | | | | | | | | | | | | | |
|---------|---------------------------------|-----|------|-----|-----|-----|------|----------------|-----|------|------|------|-----|-----|-----|------|-----|--|
| 110 | | | | | | | | US 2004-751388 | | | | | | | | | | |
| | | | - | | | | | WO 2005-US223 | | | | | | | | | | |
| WO | WO 2005068446 W: AE, AG, AL, | | | | | | | | | | | | _ | | | | | |
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| | | CN, | co, | CR, | CU, | CZ, | DE, | DK, | DM, | DZ, | EC, | EE, | EG, | ES, | FI, | GB, | GD, | |
| | | GE, | GH, | GM, | HR, | HU, | ID, | IL, | IN, | IS, | JP, | KE, | KG, | ΚP, | KR, | KZ, | LC, | |
| | | LK, | LR, | LS, | LT, | LU, | LV, | MA, | MD, | MG, | MK, | MN, | MW, | MX, | MZ, | NA, | NI, | |
| | | NO, | NZ, | OM, | PG, | PH, | PL, | PT, | RO, | RU, | SC, | SD, | SE, | SG, | SK, | SL, | SY, | |
| | | ТJ, | TM, | TN, | TR, | TT, | TZ, | UA, | UG, | US, | UΖ, | VC, | VN, | YU, | ZA, | ZM, | zw | |
| | RW: | BW, | GH, | GM, | KE, | LS, | MW, | MZ, | NA, | SD, | SL, | SZ, | TZ, | UG, | ZM, | ZW, | AM, | |
| | | AZ, | BY, | KG, | KZ, | MD, | RU, | ТJ, | TM, | AT, | BE, | BG, | CH, | CY, | CZ, | DE, | DK, | |
| | | EE, | ES, | FI, | FR, | GB, | GR, | HU, | ΙE, | IS, | IT, | LT, | LU, | MC, | NL, | PL, | PT, | |
| | | RO, | SE, | SI, | SK, | TR, | BF, | ВJ, | CF, | CG, | CI, | CM, | GA, | GN, | GQ, | GW, | ML, | |
| | | MR, | ΝE, | SN, | TD, | TG | | | | | | | | | | | | |
| PRIORIT | Y APP | LN. | INFO | . : | | | | | 1 | US 2 | 004- | 7513 | 53 | (. | A 2 | 0040 | 105 | |
| | | | | | | | | | 1 | US 2 | 004- | 7513 | 54 | | A 2 | 0040 | 105 | |
| | | | | | | | | | 1 | US 2 | 004- | 7513 | 87 | | A 2 | 0040 | 105 | |
| | | | | | | | | | 1 | US 2 | 004- | 7513 | 88 | | A 2 | 0040 | 105 | |
| | | | | | | | | | 1 | US 2 | 004- | 8226 | 39 | | A 2 | 0040 | 412 | |
| | | | | | | | | | 1 | US 2 | 004- | 8527 | 30 | | A 2 | 0040 | 524 | |
| OTHER S | OTHER SOURCE(S): | | | | | PAT | 143: | 1155 | 74 | | | | | | | | | |

GΙ

Isotopically enriched N-substituted piperazines (I) or salts thereof, comprising one or more heavy atom isotopes (Y = straight chain or branched C1-6 alkyl or C1-6 alkyl ether group wherein the carbon atoms of the alkyl group or alkyl ether group each independently comprise linked hydrogen, deuterium or fluorine atoms; Z = independently H, F, Cl, Br, iodine, an amino acid side chain, a straight chain or branched C1-6 alkyl group that may optionally contain a substituted or unsubstituted aryl group wherein the carbon atoms of the alkyl and aryl groups each independently comprise linked H or F atoms, a straight chain or branched C1-6 alkyl ether group that may optionally contain a substituted or unsubstituted aryl group (wherein the carbon atoms of the alkyl and aryl groups each independently comprise linked hydrogen or fluorine atoms), or a straight chain or branched C1-6 alkoxy group that may optionally contain a substituted or unsubstituted aryl group; wherein the carbon atoms of the alkyl and aryl groups each independently comprise linked hydrogen or fluorine atoms; wherein the N-methylpiperazine is isotopically enriched with either of 13C and/or 15N) are prepared N-substituted piperazines can be used as intermediates in the synthesis of N-substituted piperazine acetic acids

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which in turn can be used as intermediates in the synthesis of active esters of N-substituted piperazine acetic acid. The active esters of N-substituted piperazine acetic acid can be used as labeling reagents to prepare a set of isobaric labeling reagents. The set of isobaric labeling reagents can be used to label analytes such as peptides, proteins, amino acids, oligonucleotides, DNA, RNA, lipids, carbohydrates, steroids, small mols. and the like (no data). Thus, to a stirring solution of 1.18 g (11.83 mmol) N-methylpiperazine in 15 mL toluene at room temperature was added 1 g (5.91 mmol) of Et bromoacetate-1,2-13C dropwise, over a period of 15 min. The reaction mixture was then heated in an oil bath at 90° for 4 h, cooled to room temperature, filtered to remove the off-white solid to give, after workup on the combined filtrate and washings, 1.10 g (quant.) of 4-methylpiperazine-1-acetic acid Et ester-1,2-13C (II) as an off-white oil. II (1.1 g) was refluxed in water for 24 h to give 780 mg 4-methylpiperazine-1-acetic acid-1,2-13C.

IT 856188-20-0P

CN

RL: ARG (Analytical reagent use); SPN (Synthetic preparation); ANST (Analytical study); PREP (Preparation); USES (Uses)

(preparation of isotopically enriched N-substituted piperazines as isobaric labeling reagents)

RN 856188-20-0 HCAPLUS

2,5-Pyrrolidinedione, 1-[[(4-methyl-1-piperazinyl-1-15N)acetyl-2-13C-180]oxy]-, dihydrochloride (9CI) (CA INDEX NAME)

•2 HCl

IT 856188-16-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of isotopically enriched N-substituted piperazines as isobaric labeling reagents)

RN 856188-16-4 HCAPLUS

•2 HCl

RN 856188-06-2 HCAPLUS
CN 2,5-Pyrrolidinedione, 1-[[(4-methyl-1-piperazinyl)acetyl]oxy]- (9CI) (CA
INDEX NAME)

$$\begin{array}{c|c} & & & \\ & & \\ N & & \\ Me & & \\ \end{array}$$

RN 857027-09-9 HCAPLUS
CN 2-Pyrrolidinone, 1-[[(4-methyl-1-piperazinyl)acetyl]oxy]- (9CI) (CA INDEX NAME)

RN 857027-10-2 HCAPLUS

CN 1-Piperazineacetic acid, 4-methyl-, pentafluorophenyl ester (9CI) (CA INDEX NAME)

Me N
$$CH_2$$
 CH_2 F

RN 857503-00-5 HCAPLUS

CN 1-Piperazineacetic acid, 4-methyl-, pentachlorophenyl ester (9CI) (CA INDEX NAME)

RN 857503-01-6 HCAPLUS

CN 1-Piperazineacetic acid, 4-methyl-, 4-nitrophenyl ester (9CI) (CA INDEX NAME)

RN 857503-03-8 HCAPLUS

CN 1-Piperazineacetic acid, 4-methyl-, 3-nitrophenyl ester (9CI) (CA INDEX NAME)

$$N - CH_2 - C - O$$
 NO_2

L8 ANSWER 2 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2005:592129 HCAPLUS

DOCUMENT NUMBER:

143:97398

TITLE:

Preparation of active esters of N-substituted piperazine acetic acids, including isotopically

enriched versions

INVENTOR (S):

Dey, Subhakar; Pappin, Darryl J. C.; Purkayastha,

Subhasish; Pillai, Sasi; Coull, James M.

PATENT ASSIGNEE(S):

Applera Corp., USA

SOURCE:

U.S. Pat. Appl. Publ., 33 pp.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT: 6

PATENT INFORMATION:

| P. | | | | | | KIND DATE | | | APPLICATION NO. | | | | | | DATE | | | |
|--------|-------------------|------|-----|------|-----|-------------|---------|------|---------------------------------|-----|------|------|------|-----|------|-----|------|-----|
| _ | _ | | | 71 | | A1 20050707 | | | US 2004-751354 WO 2005-US223 | | | | | | | | | |
| W | | | | | | | | | | | | | | | | | | |
| | | W: | • | • | • | | | | - | | | • | | | | | | |
| | | | • | • | • | | | DE, | • | | | • | - | - | - | | | |
| | | | - | | | | | ID, | | | | | | | | | | |
| | | | - | - | | | | LV, | | | | | | | | | | |
| | | | NO, | NZ, | OM, | PG, | PH, | PL, | PT, | RO, | RU, | SC, | SD, | SE, | SG, | SK, | SL, | SY, |
| | | | TJ, | TM, | TN, | TR, | TT, | TZ, | UΑ, | ŪĠ, | US, | UΖ, | VC, | VN, | ΥU, | ZA, | ZM, | ZW |
| | | RW: | BW, | GH, | GM, | KE, | LS, | MW, | MZ, | NA, | SD, | SL, | SZ, | TZ, | UG, | ZM, | ZW, | AM, |
| | | | AZ, | BY, | KG, | ΚZ, | MD, | RU, | TJ, | TM, | AT, | BE, | BG, | CH, | CY, | CZ, | DE, | DK, |
| | | | EE, | ES, | FI, | FR, | GB, | GR, | HU, | IE, | IS, | IT, | LT, | LU, | MC, | NL, | PL, | PT, |
| | | | - | | | | | BF, | | | | | | | | | | |
| | | | MR, | NE, | SN, | TD, | TG | | | | | | | | | | | |
| PRIORI | TY | APPI | LN. | INFO | . : | | | | | 1 | US 2 | 004- | 7513 | 53 | 1 | A 2 | 0040 | 105 |
| | | | | | | | | | | 1 | US 2 | 004- | 7513 | 54 | 1 | A 2 | 0040 | 105 |
| | | | | | | | | | | 1 | US 2 | 004- | 7513 | 87 | 1 | A 2 | 0040 | 105 |
| | | | | | | | | | | 1 | US 2 | 004- | 7513 | 88 | | A 2 | 0040 | 105 |
| | | | | | | | | | | | | 004- | | | | A 2 | 0040 | 412 |
| | | | | | | | | | | 1 | US 2 | 004- | 8527 | 3.0 | | A 2 | 0040 | 524 |
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OTHER SOURCE(S):

MARPAT 143:97398

GI

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AB In some embodiments, this invention pertains to active esters of N-substituted piperazine acetic acid I (R = leaving group; X = O, S; Y = C1-C6 alkyl, C1-C6 alkyl ether; Z = H, 2H, F, Cl, Br, iodide, amino acid side chain, C1-C6 alkyl, C1-C6 alkyl ether), including isotopically enriched versions thereof. In some embodiments, this invention pertains to methods for the preparation of active esters of N-substituted piperazine acetic acid, including isotopically enriched versions thereof. For example, the isotopically labeled N-methylpiperazine II (R1 = 180H) reacted with the trifluoroacetic acid ester of N-hydroxysuccinimide to give the succinate II (R1 = OR2, R2 = succinimido).

IT 856187-87-6P 856188-06-2P 856188-16-4P 856188-20-0P

RL: IMF (Industrial manufacture); SPN (Synthetic preparation); PREP (Preparation)

(preparation of active esters of N-substituted piperazine acetic acids and their labeled derivs.)

RN 856187-87-6 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[[(4-methyl-1-piperazinyl)acetyl-180]oxy]- (9CI) (CA INDEX NAME)

RN 856188-06-2 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[[(4-methyl-1-piperazinyl)acetyl]oxy]- (9CI) (CA INDEX NAME)

N—
$$CH_2$$
— CH_2 — CH_2

RN 856188-16-4 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[[(4-methyl-1-piperazinyl)acetyl-13C2-180]oxy]-, dihydrochloride (9CI) (CA INDEX NAME)

●2 HCl

856188-20-0 HCAPLUS RN

2,5-Pyrrolidinedione, 1-[[(4-methyl-1-piperazinyl-1-15N)acetyl-2-13C-CN180]oxy]-, dihydrochloride (9CI) (CA INDEX NAME)

●2 HCl

ANSWER 3 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2005:592027 HCAPLUS

DOCUMENT NUMBER: 143:93642

TITLE: Mixtures of isobarically labeled analytes and

fragments ions derived therefrom

INVENTOR(S): Pappin, Darryl J. C.; Purkayastha, Subhasish; Coull,

James M.

PATENT ASSIGNEE(S): Applera Corp., USA

U.S. Pat. Appl. Publ., 36 pp., Cont.-in-part of U.S. Ser. No. 751,353. SOURCE:

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO. | KIND D | DATE A | APPLICATION NO. | DATE |
|----------------|-----------|-------------|-----------------|-----------------|
| | | | | |
| US 2005147985 | A1 2 | 20050707 U | JS 2004-822639 | 20040412 |
| US 2005147982 | A1 2 | 20050707 U | IS 2004-751353 | 20040105 |
| US 2005148087 | A1 2 | 20050707 U | JS 2004-852730 | 20040524 |
| WO 2005068446 | A1 2 | 20050728 W | 7O 2005-US223 | 20050105 |
| W: AE, AG, AL, | AM, AT, | AU, AZ, BA, | BB, BG, BR, BW, | BY, BZ, CA, CH, |
| CN, CO, CR, | CU, CZ, I | DE, DK, DM, | DZ, EC, EE, EG, | ES, FI, GB, GD, |
| GE, GH, GM, | HR, HU, | ID, IL, IN, | IS, JP, KE, KG, | KP, KR, KZ, LC, |
| LK, LR, LS, | LT, LU, | LV, MA, MD, | MG, MK, MN, MW, | MX, MZ, NA, NI, |

Grazier 10_765267

NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG US 2004-751353 A2 20040105 PRIORITY APPLN. INFO .: US 2004-751354 A 20040105 US 2004-751387 A 20040105 US 2004-751388 A 20040105 US 2004-822639 A2 20040412 US 2004-852730 A 20040524

OTHER SOURCE(S): MARPAT 143:93642

This invention pertains to mixts. of isobarically labeled analytes and AB fragment ions thereof.

856188-06-2P 857027-09-9P IT

> RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(mixts. of isobarically labeled analytes and fragments ions derived therefrom)

856188-06-2 HCAPLUS RN

2,5-Pyrrolidinedione, 1-[[(4-methyl-1-piperazinyl)acetyl]oxy]- (9CI) CN INDEX NAME)

857027-09-9 HCAPLUS RN

2-Pyrrolidinone, 1-[[(4-methyl-1-piperazinyl)acetyl]oxy]- (9CI) (CA INDEX CN NAME)

IT 856187-87-6P 856188-16-4P 856188-20-0P

857027-10-2P

RL: SPN (Synthetic preparation); PREP (Preparation)

(mixts. of isobarically labeled analytes and fragments ions derived therefrom)

856187-87-6 HCAPLUS RN

2,5-Pyrrolidinedione, 1-[[(4-methyl-1-piperazinyl)acetyl-180]oxy]- (9CI) CN (CA INDEX NAME)

Grazier 10_765267

L8 ANSWER 4 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2005:588426 HCAPLUS

DOCUMENT NUMBER: 143:115568

TITLE: Preparation of isotopically enriched N-substituted

piperazine-1-acetic acids

INVENTOR(S): Dey, Subhakar; Pappin, Darryl J. c.; Purkayastha,

Subhasish; Pillai, Sasi; Coull, James M.

PATENT ASSIGNEE(S): Applera Corp., USA

SOURCE: U.S. Pat. Appl. Publ., 29 pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 6

PATENT INFORMATION:

| PAT | PATENT NO. | | | | | D - | DATE | | | APPLICATION NO. | | | | | | | | |
|------------------------|--------------------------------|--------|------|-----|-------|----------|------------|------|-----|---------------------------------|------|------|-----|-----|----------|--------------|-----|--|
| | US 2005148774 WO 2005068446 | | | | | | | | | US 2004-751387 WO 2005-US223 | | | | | 20040105 | | | |
| | | AE, | AG, | AL, | AM, | AT, | AU, DE, | AZ, | BA, | BB, | BG, | BR, | BW, | BY, | BZ, | CA, | CH, | |
| | | GE, | GH, | GM, | HR, | HU, | ID, | IL, | IN, | IS, | JP, | KE, | KG, | KP, | KR, | KZ, | LC, | |
| | | NO, | NZ, | OM, | PG, | PH, | PL, TZ, | PT, | RO, | RU, | sc, | SD, | SE, | SG, | sĸ, | SL, | SY, | |
| | RW: | BW, | GH, | GM, | KE, | LS, | MW, RU, | MZ, | NA, | SD, | ŞL, | SZ, | TZ, | UG, | ZM, | ZW, | AM, | |
| | | EE, | ES, | FI, | FR, | GB, | GR, BF, | HU, | IE, | ıs, | IT, | LT, | LU, | MC, | NL, | PL, | PT, | |
| PRIORITY | λDD | MR, | NE, | SN, | TD, | | Dr, | БО, | | US 2 | | | | | | | | |
| PRIORITI | REF | LIIV . | INFO | • • | | | | | | US 2 | 004- | 7513 | 54 | | A 2 | 0040 | 105 | |
| | | | | | | | | | | US 2 US 2 | 004- | 7513 | 88 | | A 2 | 0040 0040 | 105 | |
| OTHER SOURCE(S): | | | | | MAD | ידי א כו | 142. | 1166 | | US 2 | | | | | | | | |
| OTHER SOURCE(S): GI | | | | | PIAK. | FAI | T#3: | 1133 | 00 | | | | | | | | | |

RN 856188-16-4 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[[(4-methyl-1-piperazinyl)acetyl-13C2-180]oxy]-, dihydrochloride (9CI) (CA INDEX NAME)

•2 HCl

RN 856188-20-0 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[[(4-methyl-1-piperazinyl-1-15N)acetyl-2-13C-18O]oxy]-, dihydrochloride (9CI) (CA INDEX NAME)

•2 HCl

RN 857027-10-2 HCAPLUS

Isotopically enriched N-substituted piperazine-1-acetic acids (I) or salts AB thereof, comprising one or more heavy atom isotopes [X = O, S; Y =straight chain or branched C1-6 alkyl or C1-6 alkyl ether group wherein the carbon atoms of the alkyl group or alkyl ether group each independently comprise linked hydrogen, deuterium or F atoms; Z = independently H, deuterium, F, Cl, Br, iodine, an amino acid side chain, a straight chain or branched C1-6 alkyl group that may optionally contain a substituted or unsubstituted aryl group (wherein the carbon atoms of the alkyl and aryl groups each independently comprise linked H, deuterium or F atoms), a straight chain or branched C1-6 alkyl ether group that may optionally contain a substituted or unsubstituted aryl group wherein the carbon atoms of the alkyl and aryl groups each independently comprise linked H, deuterium or F atoms, or a straight chain or branched C1-6 alkoxy group that may optionally contain a substituted or unsubstituted aryl group (wherein the carbon atoms of the alkyl and aryl groups each independently comprise linked H, deuterium or F atoms)] are prepared N-substituted piperazines can be used as intermediates in the synthesis of N-substituted piperazine acetic acids which in turn can be used as intermediates in the synthesis of active esters of N-substituted piperazine acetic acid. The active esters of N-substituted piperazine acetic acid can be used as labeling reagents to prepare a set of isobaric labeling reagents. The set of isobaric labeling reagents can be used to label analytes such as peptides, proteins, amino acids, oligonucleotides, DNA, RNA, lipids, carbohydrates, steroids, small mols. and the like. Thus, to a stirring solution of 1.18 g (11.83 mmol) N-methylpiperazine in 15 mL toluene at room temperature was added 1 g (5.91 mmol) of Et bromoacetate-1,2-13C dropwise, over a period of 15 min. The reaction mixture was then heated in an oil bath at 90° for 4 h, cooled to room temperature, filtered to remove the off-white solid to give, after workup on

the

combined filtrate and washings, 1.10 g (quant.) of 4-methylpiperazine-1-acetic acid Et ester-1,2-13C (II) as an off-white oil. II (1.1 g) was refluxed in water for 24 h to give 780 mg 4-methylpiperazine-1-acetic acid-1,2-13C.

IT 856188-20-0P

RL: ARG (Analytical reagent use); SPN (Synthetic preparation); ANST (Analytical study); PREP (Preparation); USES (Uses)

(preparation of isotopically enriched N-substituted piperazine-1-acetic acids as isobaric labeling reagents)

RN 856188-20-0 HCAPLUS

2,5-Pyrrolidinedione, 1-[[(4-methyl-1-piperazinyl-1-15N)acetyl-2-13C-180]oxy]-, dihydrochloride (9CI) (CA INDEX NAME)

•2 HCl

Grazier 10_765267

856188-16-4P IT

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of isotopically enriched N-substituted piperazine-1-acetic acids as isobaric labeling reagents)

856188-16-4 HCAPLUS RN

2,5-Pyrrolidinedione, 1-[[(4-methyl-1-piperazinyl)acetyl-13C2-180]oxy]-, CN dihydrochloride (9CI) (CA INDEX NAME)

●2 HCl

856187-87-6P 856188-06-2P 857027-09-9P IT

857027-10-2P 857503-00-5P 857503-01-6P

857503-03-8P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of isotopically enriched N-substituted piperazine-1-acetic

acids as isobaric labeling reagents)

856187-87-6 HCAPLUS RN

2,5-Pyrrolidinedione, 1-[[(4-methyl-1-piperazinyl)acetyl-180]oxy]- (9CI) CN (CA INDEX NAME)

856188-06-2 HCAPLUS RN

2,5-Pyrrolidinedione, 1-[[(4-methyl-1-piperazinyl)acetyl]oxy]- (9CI) (CA CN INDEX NAME)

857027-09-9 HCAPLUS RN

2-Pyrrolidinone, 1-[[(4-methyl-1-piperazinyl)acetyl]oxy]- (9CI) (CA INDEX CN NAME)

RN 857027-10-2 HCAPLUS

CN 1-Piperazineacetic acid, 4-methyl-, pentafluorophenyl ester (9CI) (CA INDEX NAME)

Me
$$N \longrightarrow CH_2 - C \longrightarrow F$$

RN 857503-00-5 HCAPLUS

CN 1-Piperazineacetic acid, 4-methyl-, pentachlorophenyl ester (9CI) (CA INDEX NAME)

Me
$$N \longrightarrow CH_2 \longrightarrow C1$$
 $C1$ $C1$ $C1$ $C1$

RN 857503-01-6 HCAPLUS

CN 1-Piperazineacetic acid, 4-methyl-, 4-nitrophenyl ester (9CI) (CA INDEX NAME)

RN 857503-03-8 HCAPLUS

CN 1-Piperazineacetic acid, 4-methyl-, 3-nitrophenyl ester (9CI) (CA INDEX NAME)

$$N - CH_2 - C - O - NO_2$$

L8 ANSWER 5 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2005:588349 HCAPLUS

DOCUMENT NUMBER: 143:112150

TITLE: Isobarically labeled analytes and fragment ions

derived therefrom

INVENTOR(S): Pappin, Darryl J. C.; Purkayastha, Subhasish; Coull,

James M.

PATENT ASSIGNEE(S): Applera Corporation, USA

SOURCE: U.S. Pat. Appl. Publ., 88 pp., Cont.-in-part of U.S.

Ser. No. 822,639. CODEN: USXXCO

DOCUMENT TYPE: Patent

LANGUAGE: English

FAMILY ACC. NUM. COUNT: 6

PATENT INFORMATION:

| PATENT NO. | | APPLICATION NO. | DATE | | | | |
|------------------------|-------------------|---|-----------------|--|--|--|--|
| | | *************************************** | 20040524 | | | | |
| US 2005148087 | | | | | | | |
| US 2005147982 | | US 2004-751353 | | | | | |
| US 2005147985 | A1 20050707 | US 2004-822639 | 20040412 | | | | |
| WO 2005068446 | A1 20050728 | WO 2005-US223 | 20050105 | | | | |
| W: AE, AG, AL | , AM, AT, AU, AZ, | BA, BB, BG, BR, BW, | BY, BZ, CA, CH, | | | | |
| CN, CO, CR | , CU, CZ, DE, DK, | DM, DZ, EC, EE, EG, | ES, FI, GB, GD, | | | | |
| | | IN, IS, JP, KE, KG, | | | | | |
| | | MD, MG, MK, MN, MW, | | | | | |
| | | RO, RU, SC, SD, SE, | | | | | |
| | | UG, US, UZ, VC, VN, | | | | | |
| | | NA, SD, SL, SZ, TZ, | | | | | |
| | | TM, AT, BE, BG, CH, | | | | | |
| | | IE, IS, IT, LT, LU, | | | | | |
| | | | | | | | |
| • | | CF, CG, CI, CM, GA, | GN, GQ, GW, ML, | | | | |
| MR, NE, SN | , TD, TG | | | | | | |
| PRIORITY APPLN. INFO.: | | US 2004-751353 | | | | | |
| | | US 2004-822639 | A2 20040412 | | | | |
| | | US 2004-751354 | A 20040105 | | | | |
| | | US 2004-751387 | A 20040105 | | | | |
| | | US 2004-751388 | A 20040105 | | | | |
| | | US 2004-852730 | A 20040524 | | | | |
| | | | | | | | |

OTHER SOURCE(S): MARPAT 143:112150

AB This invention pertains to isobarically labeled analytes and fragment ions thereof.

IT 857027-09-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(isobarically labeled analytes and fragment ions derived therefrom)

RN 857027-09-9 HCAPLUS

CN 2-Pyrrolidinone, 1-[[(4-methyl-1-piperazinyl)acetyl]oxy]- (9CI) (CA INDEX NAME)

IT 741683-79-4P 856187-87-6P 856188-06-2P

857027-10-2P

RL: SPN (Synthetic preparation); PREP (Preparation)

(isobarically labeled analytes and fragment ions derived therefrom)

741683-79-4 HCAPLUS RN

2,5-Pyrrolidinedione, 1-[(1-piperidinylacetyl)oxy]- (9CI) (CA INDEX NAME) CN

856187-87-6 HCAPLUS RN

2,5-Pyrrolidinedione, 1-[[(4-methyl-1-piperazinyl)acetyl-180]oxy]- (9CI) CN (CA INDEX NAME)

856188-06-2 HCAPLUS RN

2,5-Pyrrolidinedione, 1-[[(4-methyl-1-piperazinyl)acetyl]oxy]- (9CI) CNINDEX NAME)

RN857027-10-2 HCAPLUS

1-Piperazineacetic acid, 4-methyl-, pentafluorophenyl ester (9CI) (CA CNINDEX NAME)

ANSWER 6 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

2005:588336 HCAPLUS ACCESSION NUMBER:

143:93635 DOCUMENT NUMBER:

TITLE: Mixtures of isobarically labeled analytes and

fragments ions derived therefrom

Pappin, Darryl J. C.; Purkayastha, Subhasish; Coull, INVENTOR(S):

James M.

Applera Corporation, USA PATENT ASSIGNEE(S): SOURCE: U.S. Pat. Appl. Publ., 29 pp.

CODEN: USXXCO

DOCUMENT TYPE: Patent English LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PA | PATENT NO. | | | KINI |) | DATE | | APPLICATION NO. | | | | | | DATE | | | | |
|---------|---------------|------|------|-------------|-------------|------|------|-----------------|----------------|-------|------|------|------|----------|----------|-------|-----|--|
| | | | | | - · | - | | | | | | | | | - | | | |
| US | 2005 | 1479 | B2 | | A1 | | 2005 | 0707 | US 2004-751353 | | | | | | 20040105 | | | |
| US | 2005 | 1479 | 85 | | A1 | | 2005 | 0707 | US 2004-822639 | | | | | | 20040412 | | | |
| US | US 2005148087 | | | | A1 20050707 | | | US 2004-852730 | | | | | | 20040524 | | | | |
| WO | WO 2005068446 | | | A1 20050728 | | | | WO 2005-US223 | | | | | | 20050105 | | | | |
| | W: | ΑE, | AG, | AL, | AM, | AT, | AU, | AZ, | BA, | BB, | BG, | BR, | BW, | BY, | BZ, | CA, | CH, | |
| | | CN, | CO, | CR, | CU, | CZ, | DE, | DK, | DM, | DZ, | EC, | EE, | EG, | ES, | FI, | GB, | GD, | |
| | | GE, | GH, | GM, | HR, | HU, | ID, | IL, | IN, | IS, | JP, | KE, | KG, | ΚP, | KR, | ΚŻ, | LC, | |
| | | LK, | LR, | LS, | LT, | LU, | LV, | MA, | MD, | MG, | MK, | MN, | MW, | MX, | MZ, | NΑ, | NΙ, | |
| | | NO, | NZ, | OM, | PG, | PH, | PL, | PT, | RO, | RU, | SC, | SD, | SE, | SG, | SK, | SL, | SY, | |
| | | ТJ, | TM, | TN, | TR, | TT, | TZ, | UA, | UG, | US, | UΖ, | VC, | VN, | YU, | ZA, | ZM, | zw | |
| | RW: | BW, | GH, | GM, | KE, | LS, | MW, | ΜZ, | NA, | SD, | SL, | SZ, | TZ, | ŪĠ, | ZM, | ZW, | AM, | |
| | | AZ, | BY, | KG, | ΚZ, | MD, | RU, | TJ, | TM, | AT, | BE, | BG, | CH, | CY, | CZ, | DE, | DK, | |
| | | EE, | ES, | FI, | FR, | GB, | GR, | HU, | ΙE, | IS, | IT, | LT, | LU, | MC, | NL, | ΡL, | PT, | |
| | | RO, | SE, | SI, | SK, | TR, | BF, | ВJ, | CF, | CG, | CI, | CM, | GA, | GN, | GQ, | GW, | ML, | |
| | | MR, | NE, | SN, | TD, | TG | | | | | | | | | | | | |
| PRIORIT | Y APP | LN. | INFO | . : | | | | | 1 | US 2 | 004- | 7513 | 53 | | A2 2 | 0040 | 105 | |
| | | | | | | | | | 1 | US 2 | 004- | 7513 | 54 | | A 2 | 0040 | 105 | |
| | | | | | | | | | 1 | US 2 | 004- | 7513 | 87 | | A 2 | 0040 | 105 | |
| | | | | | | | | | 1 | US 2 | 004- | 7513 | 88 | | A 2 | 0040 | 105 | |
| | | | | | | | | | 7 | US 2 | 004- | 8226 | 39 | , | A2 2 | 0040 | 412 | |
| | | | | | | | | | 1 | US 2 | 004- | 8527 | 30 | | A 2 | 0040 | 524 | |
| | is in | | - | - | | | | s. o | f is | obar. | ical | ly l | abel | ed a | naly | tes . | and | |

fragment ions thereof.

856188-06-2P 857027-09-9P IT

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(mixts. of isobarically labeled analytes and fragments ions derived therefrom)

856188-06-2 HCAPLUS RN

2,5-Pyrrolidinedione, 1-[[(4-methyl-1-piperazinyl)acetyl]oxy]- (9CI) CNINDEX NAME)

$$\begin{array}{c|c} & & & \\ & & \\ N & & \\ N & & \\ \end{array}$$

RN 857027-09-9 HCAPLUS

CN 2-Pyrrolidinone, 1-[[(4-methyl-1-piperazinyl)acetyl]oxy]- (9CI) (CA INDEX NAME)

IT 856187-87-6P 857027-10-2P

RL: SPN (Synthetic preparation); PREP (Preparation)
 (mixts. of isobarically labeled analytes and fragments ions derived
 therefrom)

RN 856187-87-6 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[[(4-methyl-1-piperazinyl)acetyl-180]oxy]- (9CI) (CA INDEX NAME)

RN 857027-10-2 HCAPLUS

CN 1-Piperazineacetic acid, 4-methyl-, pentafluorophenyl ester (9CI) (CA INDEX NAME)

L8 ANSWER 7 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 2005:412987 HCAPLUS

Grazier 10_765267

DOCUMENT NUMBER: 144:186804

TITLE: Analysis of cell membrane aminophospholipids as

isotope-tagged derivatives

AUTHOR(S): Zemski Berry, Karin A.; Murphy, Robert C.

CORPORATE SOURCE: Department of Pharmacology, University of Colorado

Health Sciences Center, Aurora, CO, 80045, USA

SOURCE: Journal of Lipid Research (2005), 46(5), 1038-1046

CODEN: JLPRAW; ISSN: 0022-2275

PUBLISHER: American Society for Biochemistry and Molecular

Biology, Inc.

DOCUMENT TYPE: Journal LANGUAGE: English

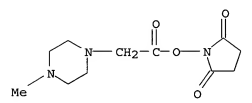
AB Glycerophosphoethanolamine (GPEtn) and glycerophosphoserine (GPSer) lipids were reacted with a multiplexed set of differentially isotopically enriched N-methylpiperazine acetic acid N-hydroxysuccinimide ester reagents, which place isobaric mass labels at a primary amino group. resulting derivitized aminophospholipids were isobaric and chromatog. indistinguishable but yielded pos. reporter ions (m/z 114 or 117) after collisional activation that could be used to identify and quantify individual members of the multiplex set. The chromatog. and mass spectrometric response of N-methylpiperazine amide-tagged aminophospholipids was probed using glycerophosphoethanolamine and glycerophosphoserine lipid stds. The [M+H]+ of each tagged aminophospholipid shifted 144 Da, and during collision-induced dissociation the major fragmentation ion was either m/z 114 or 117. This mode of detecting aminophospholipids was useful for an unbiased anal. of plasmalogen GPEtn lipids. Mol. species information on the esterified fatty acyl substituents was obtained by collisional activation of the [M-H] - ions. The isotope-tagged reagents were used to assess changes in the distribution of GPEtn lipids after exposure of liposomes made from phospholipids extracted from RAW 264.7 cells to Cu2+/H2O2 to illustrate the ability of these reagents to aid in the mass spectrometric identification of aminophospholipid changes that occur during biol. stimuli.

IT 856188-06-2

RL: RCT (Reactant); RACT (Reactant or reagent)
(preparation and mass spectrometric anal. of cell membrane
aminophospholipids as isotope-tagged derivs.)

RN 856188-06-2 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[[(4-methyl-1-piperazinyl)acetyl]oxy]- (9CI) (CAINDEX NAME)



REFERENCE COUNT: 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 8 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2005:271531 HCAPLUS

DOCUMENT NUMBER: 142:492702

TITLE: Coordination Behavior toward Copper(II) and Zinc(II)

Ions of Three Ligands Joining 3-Hydroxy-2-pyridinone

and Polyaza Fragments

Grazier 10_765267

AUTHOR(S): Ambrosi, Gianluca; Formica, Mauro; Fusi, Vieri;

Giorgi, Luca; Guerri, Annalisa; Lucarini, Simone; Micheloni, Mauro; Paoli, Paola; Rossi, Patrizia;

Zappia, Giovanni

CORPORATE SOURCE: Institute of Chemical Sciences and Institute of

Pharmaceutical Chemistry, University of Urbino,

Urbino, I-61029, Italy

SOURCE: Inorganic Chemistry (2005), 44(9), 3249-3260

CODEN: INOCAJ; ISSN: 0020-1669

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

GI

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The synthesis and characterization of new polydentate ligand 2-(N),2'-(N')-bis[2-(3-hydroxy-2-oxo-2H-pyridin-1-yl)acetamido]-1(N'),2(N),2'(N')-trimethyl-2,2'-diaminodiethylamine, I (L3), is reported. The coordination properties of L3 and of two analogous macrocyclic ligands II [Z = H2, (L1) and Z = O (L2)] toward Cu(II) and Zn(II) metal ions are reported. All three ligands show the 3-hydroxy-2(1H)-pyridinone (HPO) groups attached as sidearms to a polyaza fragment, which is a macrocyclic framework in the case of L1 and L2 while it is an open chain in the case of L3. The role of the polyaza fragments in preorganizing the two sidearms was studied. The basicity of L3 and the binding properties of L1-L3 were determined by potentiometric measurements in aqueous solution (298.1 ±

0.1 K, I = 0.15 mol dm-3). UV-visible spectra as well 1H and 13C NMR expts. were used to understand the role of the HPO and of the polyaza fragments in the stabilization of the cations. While L1 forms stable mono- and dinuclear complexes, L2 and L3 can form only mononuclear species with each of the metal ions studied. In the main mononuclear species of L2 and L3, the two HPO moieties stabilize the M(II) in a square-planar geometry due to the two O atoms of each HPO. The coordination sphere of the metal is completed by adding a secondary ligand such as H2O mols. in the case of Cu(II) systems or OH- in the Zn(II) systems. These results are confirmed by the crystal structures of the [CuH-1L2]+ and [CuH-1L3]+ species reported herein. Two conformations of L1 can be hypothesized in the formation of the dinuclear species, as suggested by NMR expts. on the [ZnH-2L1] species, which shows two conformers slowly interchanging on the NMR time scale, one of which is more insol.

IT 852159-56-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(for preparation of polyaza hydroxypyridinone-containing ligand)

RN 852159-56-9 HCAPLUS

CN 1(2H)-Pyridineacetic acid, 2-oxo-3-(phenylmethoxy)-, 4-nitrophenyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT: 48 THERE ARE 48 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 9 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:824132 HCAPLUS

DOCUMENT NUMBER: 141:310231 TITLE: Mass labels

INVENTOR(S): Hamon, Christian; Kuhn, Karsten; Thompson, Andrew;

Reuschling, Dieter; Schaefer, Juergen

PATENT ASSIGNEE(S): Xzillion G.m.b.H. & Co. K.-G., Germany; Proteome

Sciences PLC

SOURCE: PCT Int. Appl., 63 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| | PATENT NO. | | | | | | | | APPLICATION NO. | | | | | | DATE | | | |
|---------------|--------------------------------|-----|-----|-------------|-----|------|----------------|------|-----------------|------|-------|------|----------|------------|------|------|-----|--|
| WO | WO 2004086050 WO 2004086050 | | | A2 20041007 | | | WO 2004-GB1167 | | | | | | 20040318 | | | | | |
| | W: | ΑE, | AG, | AL, | AM, | ΑT, | AU, | ΑZ, | BA, | BB, | BG, | BR, | BW, | BY, | ΒZ, | CA, | CH, | |
| | | CN, | CO, | CR, | CU, | CZ, | DE, | DK, | DM, | DZ, | EC, | EE, | EG, | ES, | FI, | GB, | GD, | |
| | | GE, | GH, | GM, | HR, | HU, | ID, | IL, | IN, | IS, | JP, | ΚE, | KG, | KP, | KR, | ΚZ, | LC, | |
| | | LK, | LR, | LS, | LT, | LU, | LV, | MA, | MD, | MG, | MK, | MN, | MW, | MX, | MZ, | NΑ, | NI, | |
| | | NO, | ΝZ, | OM, | PG, | PH, | PL, | PT, | RO, | RU, | SC, | SD, | SE, | SG, | SK, | SL, | SY, | |
| | | ТJ, | TM, | TN, | TR, | TT, | TZ, | UA, | UG, | US, | UZ, | VC, | VN, | YU, | ZA, | ZM, | ZW | |
| | RW: | BW, | GH, | GM, | KE, | LS, | MW, | MZ, | SD, | SL, | SZ, | TZ, | UG, | ZM, | ZW, | AM, | ΑZ, | |
| | | | • | | • | | | TM, | - | - | - | - | | | | | - | |
| | | ES, | FI, | FR, | GB, | GR, | HU, | ΙE, | IT, | LU, | MC, | NL, | PL, | PT, | RO, | SE, | SI, | |
| | | SK, | TR, | BF, | ВJ, | CF, | CG, | CI, | CM, | GΑ, | GN, | GQ, | GW, | ML, | MR, | ΝE, | SN, | |
| | | TD, | TG | | | | | | | | | | | | | | | |
| CA | CA 2520297 | | | | AA | | 2004 | 1007 | • | CA 2 | 004- | 2520 | 297 | | 2 | 0040 | 318 | |
| EP 1606623 | | | | A2 | | 2005 | 1221 |] | EP 2 | 004- | 7215 | 65 | | 2 | 0040 | 318 | | |
| | R: | ΑT, | BE, | CH, | DE, | DK, | ES, | FR, | GB, | GR, | IT, | LI, | LU, | ΝL, | SE, | MC, | PT, | |
| | | ΙE, | SI, | LT, | LV, | FI, | RO, | MK, | CY, | AL, | TR, | BG, | CZ, | EE, | ΗU, | PL, | SK | |
| NO 2005004684 | | | | | Α | | 2005 | 1012 | NO 2005-4684 | | | | | 20051012 | | | | |
| PRIORIT | PRIORITY APPLN. INFO.: | | | | | | | | GB 2003-6756 | | | | | A 20030324 | | | | |
| | | | | | | | | | 1 | WO 2 | 004-0 | GB11 | 67 | ı | v 2 | 0040 | 318 | |

- AB Provided is a method for characterizing a mol. by mass spectrometry, which mol. comprises one or more free amino groups, which method comprises: (a) reacting one or more free amino groups in the mol. with a mass tag reagent comprising a reactive functionality capable of reacting with an amino group, and a tertiary amino group linked to the reactive functionality; and (b) characterizing the mol. by mass spectrometry.
- IT 741683-76-1P 741683-79-4P 768385-34-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(mass labels)

RN 741683-76-1 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[(4-morpholinylacetyl)oxy]- (9CI) (CA INDEX NAME)

RN 741683-79-4 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[(1-piperidinylacetyl)oxy]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c}
 & \circ & \circ \\
 & \circ & \circ \\
 & \circ & \circ \\
 & \circ & \circ
\end{array}$$

RN 768385-34-8 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[[(2,6-dimethyl-1-piperidinyl)acetyl]oxy]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} Me & O & O \\ \hline N - CH_2 - C - O - N \\ \hline Me & O \\ \end{array}$$

L8 ANSWER 10 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:681717 HCAPLUS

DOCUMENT NUMBER: 141:202794

TITLE: Methods, mixtures, kits and compositions pertaining to

analyte determination

INVENTOR(S): Pappin, Darryl J. C.; Bartlet-Jones, Michael

PATENT ASSIGNEE(S): Applera Corporation, USA SOURCE: PCT Int. Appl., 105 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | KIND DATE | APPLICATION NO. | DATE | | |
|----------------|----------------|----------------------------|-------------|--|--|
| | | | | | |
| WO 2004070352 | A2 2004083 | L9 WO 2004-US2077 | 20040127 | | |
| W: AE, AG, AL, | AM, AT, AU, AZ | Z, BA, BB, BG, BR, BW, BY, | BZ, CA, CH, | | |
| CN, CO, CR, | CU, CZ, DE, DI | K, DM, DZ, EC, EE, EG, ES, | FI, GB, GD, | | |
| GE, GH, GM, | HR, HU, ID, II | I, IN, IS, JP, KE, KG, KP, | KR, KZ, LC, | | |

Grazier 10 765267

LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI RW: BW, GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG CA 2488584 AA 20040819 CA 2004-2488584 20040127 US 2004-765264 US 2004219685 Α1 20041104 20040127 US 2004-765267 20041104 20040127 US 2004220412 Α1 US 2004-765458 US 2004219686 Α1 20041104 20040127 EP 1588145 20051026 EP 2004-705571 A2 20040127 AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, R: IE, SI, LT, LV, FI, RO, MK, CY, AL, TR, BG, CZ, EE, HU, SK US 2003-443612P PRIORITY APPLN. INFO.: P 20030130 W 20040127 WO 2004-US2077 This invention pertains to methods, mixts., kits and/or compns. for the AB determination of analytes by mass anal. using unique labeling reagents or sets of unique labeling reagents. The labeling reagents can be isomeric or isobaric and can be used to produce mixts. suitable for multiplex anal. of the labeled analytes. 741683-76-1P 741683-77-2P 741683-78-3P IT 741683-79-4P 741683-80-7P 741683-86-3P 741683-93-2P RL: SPN (Synthetic preparation); PREP (Preparation) (methods, mixts., kits and compns. pertaining to analyte determination) RN741683-76-1 HCAPLUS 2,5-Pyrrolidinedione, 1-[(4-morpholinylacetyl)oxy]- (9CI) (CA INDEX NAME) CN

RN 741683-77-2 HCAPLUS
CN 2,5-Pyrrolidinedione, 1-[(4-morpholinylacetyl-1-13C)oxy]- (9CI) (CA INDEX NAME)

RN 741683-78-3 HCAPLUS
CN 2,5-Pyrrolidinedione, 1-[(4-morpholinylacetyl-2-13C)oxy]- (9CI) (CA INDEX NAME)

RN 741683-79-4 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[(1-piperidinylacetyl)oxy]- (9CI) (CA INDEX NAME)

RN 741683-80-7 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[(1-piperazinylacetyl)oxy]- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c}
 & O & O & O \\
 & O & O &$$

RN 741683-86-3 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[(1-piperidinylacetyl-1-13C)oxy]- (9CI) (CA INDEX NAME)

RN 741683-93-2 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[(1-piperidinylacetyl-2-13C)oxy]- (9CI) (CA INDEX NAME)

L8 ANSWER 11 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:984421 HCAPLUS

DOCUMENT NUMBER: 140:163851

TITLE: New ligand bearing preorganized binding side-arms

interacting with ammonium cations: synthesis, conformational studies and crystal structure

AUTHOR(S): Formica, Mauro; Fusi, Vieri; Giorgi, Luca; Guerri,

Annalisa; Sucarini, Simone; Micheloni, Mauro; Paoli, Paola; Pontellini, Roberto; Rossi, Patrizia; Tarzia,

Giorgio; Zappia, Giovanni

CORPORATE SOURCE: Institute of Chemical Sciences, University of Urbino,

Urbino, I-61029, Italy

SOURCE: New Journal of Chemistry (2003), 27(11), 1575-1583

CODEN: NJCHE5; ISSN: 1144-0546

PUBLISHER: Royal Society of Chemistry

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 140:163851

The synthesis and characterization of the new tetraazamacrocycle 4-(N)-bis[2-(3-hydroxy-2-oxo-2H-pyridin-1-yl)acetamido]-1,7-dimethyl-1,4,7,10-tetraazacyclododecane (I) is reported. I shows two 3-(hydroxy)-1-(carbonylmethylen)-2-(1H)-pyridinone moieties as side-arms of a tetra-aza-macrocyclic base. The key coupling of side-arms was studied and the most significant results were obtained by activating the 3-(benzyloxy)-1-(carboxymethyl)-2-(1H)-pyridinone as a pentafluorophenol ester. The acid-base properties of I and its capability to interact with simple ammonium cations were investigated by potentiometric measurements in aqueous solution (298.1±0.1 K, I = 0.15 mol dm-3). Protonated species of I can bind NH4+ or primary ammonium cations such as MeNH3+ which are not bound in aqueous solution 1H and 13C NMR spectra showed the existence in solution

of two conformers on the NMR time scale due to the rotational restriction of the two N-C=O groups. The activation parameters were determined by dynamic variable-temperature NMR anal. Mol. dynamics calcns. gave results in agreement with the exptl. data for both conformation and ammonium-binding studies, underlining that the transformation of the two secondary amines of the macrocyclic base to amide functions, forces the side-arms to remain fixed in position, almost face to face and thus to be preorganized to interact with other species. The crystal structure of the [HL]Cl.8H2O species shows the high number of preorganized hydrogen bond sites capable, in this case, of interacting directly with five H2O mols.

IT 654637-03-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(coupling reaction of; preparation and potentiometric titration of multidentate

azacrown ether ligand bearing preorganized binding side-arms interacting with ammonium cations)

RN 654637-03-3 HCAPLUS

CN 1(2H)-Pyridineacetic acid, 2-oxo-3-(phenylmethoxy)-, pentafluorophenyl ester (9CI) (CA INDEX NAME)

$$Ph-CH_2-O$$

$$N-CH_2-C-O$$

$$F$$

$$F$$

THERE ARE 67 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 12 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN $^{\text{L8}}$

67

ACCESSION NUMBER: 2003:930975 HCAPLUS

DOCUMENT NUMBER: 139:395945

Preparation of quinazolinylmethyl urea derivatives as TITLE:

fungal efflux pump inhibitors

INVENTOR(S): Watkins, Will J.; Lemoine, Remy; Cho, Aesop; Palme,

Monica

USA PATENT ASSIGNEE(S):

U.S. Pat. Appl. Publ., 109 pp., Cont.-in-part of U.S. SOURCE:

Ser. No. 906,864.

CODEN: USXXCO

Patent DOCUMENT TYPE: English LANGUAGE:

FAMILY ACC. NUM. COUNT: 3

PATENT INFORMATION:

| PA' | TENT 1 | NO. | | | KINI |) | DATE | | | APPL | ICAT: | ION I | . 01 | | D | ATE | | |
|---------|--------|------|------|-----|-------------|-------------|-------------------------|------|-----|----------------|-------|-------|----------|-----|------------|-------|-----|--|
| IIS | 2003: | 2203 | 38 | | Δ1 | - | 2003 | 1127 | , | US 2 | 002- | 2430' | 74 | | 20 | 00209 | 912 | |
| | 6596 | | | | B1 20030722 | | | | | US 2001-906864 | | | | | 20010716 | | | |
| | 2003 | | 97 | | A1 20031211 | | | | | US 2002-334755 | | | | | | | | |
| | | | | | | B2 20040210 | | | | | | | | • | | | | |
| | •• ••• | | | | | | 20040325 WO 2003-US5184 | | | | | | 20030221 | | | | | |
| | W: | | | | | | | | | | BG, | | | | | | | |
| | | • | | • | • | • | • | • | • | • | EE, | • | • | | • | • | - | |
| | | • | • | • | - | | | • | | | KG, | | | | | | | |
| | | LS, | LT, | LU, | LV, | MA, | MD, | MG, | MK, | MN, | MW, | MX, | MZ, | NO, | NZ, | OM, | PH, | |
| | | PL, | PT, | RO, | RU, | SC, | SD, | SE, | SG, | SK, | SL, | TJ, | TM, | TN, | TR, | TT, | TZ, | |
| | | UA, | ŪĠ, | UZ, | VC, | VN, | YU, | ZA, | ZM, | ZW | | | | | | | | |
| | RW: | GH, | GM, | KE, | LS, | MW, | MZ, | SD, | SL, | SZ, | TZ, | ŪĠ, | ZM, | ZW, | AM, | ΑZ, | BY, | |
| | | KG, | ΚZ, | MD, | RU, | ТJ, | TM, | AT, | BE, | BG, | CH, | CY, | CZ, | DE, | DK, | EE, | ES, | |
| | | FI, | FR, | GB, | GR, | HU, | ΙE, | IT, | LU, | MC, | NL, | PT, | SE, | SI, | SK, | TR, | BF, | |
| | | ВJ, | CF, | CG, | CI, | CM, | GΑ, | GN, | GQ, | GW, | ML, | MR, | NE, | SN, | TD, | TG | | |
| AU | 2003 | 2153 | 43 | | A1 | | 2004 | 0430 | | AU 2 | 003- | 2153 | 43 | | 2 | 0030 | 221 | |
| PRIORIT | Y APP | LN. | INFO | . : | | | | | | US 2 | 001- | 9068 | 64 | 7 | A2 2 | 0010 | 716 | |
| | | | | | | | | | • | US 2 | 002- | 2430 | 74 | Ž | A2 2 | 0020 | 912 | |
| | | | | | | | | | | US 2 | 002- | 3347 | 55 | Ž | A 2 | 0021 | 230 | |
| | | | | | | | | | | WO 2 | 003- | US51 | 84 | 1 | <i>N</i> 2 | 0030 | 221 | |

OTHER SOURCE(S): MARPAT 139:395945

GI

This invention relates to compds. of formula I [A1-A6 = C, N; R1 = H, alkyl, cycloalkyl, CH2-cycloalkyl, etc.; R2 = alkyl; R3-R12 = H, alkyl, CF3, alkoxy, halo, OH, CN, etc.] that are efflux pump inhibitors and therefore are useful as potentiators of anti-fungal agents for the treatment of infections caused by fungi that employ an efflux pump resistance mechanism. Thus, II was prepared and showed a reduced MIC value against Candida albicans in the presence of fluconazole.

IT 626245-59-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of quinazolinylmethyl urea derivs. as fungal efflux pump inhibitors)

RN 626245-59-8 HCAPLUS

CN Urea, N-(2,4-dimethoxyphenyl)-N-[1-[3-[4-[2-[(2,5-dioxo-1-pyrrolidinyl)oxy]-2-oxoethyl]-1-piperazinyl]-3,4-dihydro-4-oxo-2-quinazolinyl]ethyl]-N'-(4-fluorophenyl)- (9CI) (CA INDEX NAME)

L8 ANSWER 13 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1999:439195 HCAPLUS

DOCUMENT NUMBER: 131:184754

TITLE: Synthesis of 3-Hydroxy-2-pyridinone Derivatives of

4-tert-Butylcalix[4] arenes: A New Class of Selective

Extractants of Actinide(IV) Ions

AUTHOR(S): Lambert, Timothy N.; Dasaradhi, Lakkaraju; Huber,

Vincent J.; Gopalan, Aravamudan S.

CORPORATE SOURCE: Department of Chemistry and Biochemistry, New Mexico

State University, Las Cruces, NM, 88003-8001, USA

SOURCE: Journal of Organic Chemistry (1999), 64(16), 6097-6101

CODEN: JOCEAH; ISSN: 0022-3263

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 131:184754

GΙ

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

AB Two hydroxypyridinone calixarene derivs. I (n = 3; X = NH; X1 = 0; n = 2; X = 0; X1 = H2) have been developed as a new class of extractants for actinides. I proved efficient for extracting Th(IV) and Fe(III) and selective for Th(IV) under competitive conditions.

IT 95215-73-9

RL: RCT (Reactant); RACT (Reactant or reagent)
(preparation of hydroxypyridinone derivs. of calixarenes as selective
extractants of actinide (IV) ions)

RN 95215-73-9 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[[[2-oxo-3-(phenylmethoxy)-1(2H)-pyridinyl]acetyl]oxy]- (9CI) (CA INDEX NAME)

THERE ARE 41 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 41 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 14 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

1998:402422 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 129:81670

Preparation of therapeutic antioxidants for TITLE:

Alzheimer's disease

Tilbrook, Gary Stuart; Hider, Robert Charles; Moridani, Majid Yousefi INVENTOR(S):

Cenes Ltd., UK; Tilbrook, Gary Stuart; Hider, Robert PATENT ASSIGNEE(S):

Charles; Moridani, Majid Yousefi

PCT Int. Appl., 31 pp. SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE: Patent English LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PA | CENT 1 | NO. | | | KINI |) | DATE | | 7 | PPL | ICAT: | I NO | 10. | | D | ATE | |
|---------|------------|-------|------|-----|-----------|-----|-------|------|-----|------|-------|-------|-----|-----|------|-------|-----|
| | - - | | | | | - | | | - | | | | | | | | |
| WO | 9825 | 905 | | | A2 | | 19980 | 0618 | W | 10 1 | 997-0 | 3B33(|)6 | | 1 | 99712 | 210 |
| WO | 9825 | 905 | | | A3 | | 1998 | 1029 | | | | | | | | | |
| | W: | AL, | AM, | AT, | AU, | ΑZ, | ВA, | BB, | BG, | BR, | BY, | CA, | CH, | CN, | CU, | CZ, | DE, |
| | | DK, | EE, | ES, | FI, | GB, | GE, | GH, | GM, | HU, | ID, | ΙL, | IS, | JP, | KE, | KG, | ΚP, |
| | | KR, | KZ, | LC, | LK, | LR, | LS, | LT, | LU, | LV, | MD, | MG, | MK, | MN, | MW, | MX, | NO, |
| | | NZ, | PL, | PT, | RO, | RU, | SD, | SE, | SG, | SI, | SK, | SL, | TJ, | TM, | TR, | TT, | UA, |
| | | UG, | US, | UΖ, | VN, | YU, | ZW, | AM, | AZ, | BY, | KG, | ΚZ, | MD, | RU, | ТJ, | TM | |
| | RW: | GH, | GM, | KE, | LS, | MW, | SD, | SZ, | UG, | ZW, | ΑT, | BE, | CH, | DE, | DK, | ES, | FI, |
| | | FR, | GB, | GR, | ΙE, | IT, | LU, | MC, | NL, | PT, | SE, | BF, | ΒJ, | CF, | CG, | CI, | CM, |
| | | GA, | GN, | ML, | MR, | NE, | SN, | TD, | TG | | | | | | | | |
| AU | 9878 | 471 | | | A1 | | 1998 | 0703 | P | \U 1 | 998-1 | 7847 | 1 | | 1: | 99712 | 210 |
| PRIORIT | APP | LN. | INFO | . : | | | | | G | B 1 | 996-2 | 25638 | 3 | 7 | A 1: | 99612 | 210 |
| | | | | | | | | | V | VO 1 | 997-0 | 3B330 | 06 | 7 | N 1 | 99712 | 210 |
| OTHER S | איווערב | (9) . | | | MARI | ጥፈር | 129 - | R167 | 0 | | | | | | | | |

OTHER SOURCE(S): MARPAT 129:81670

GI

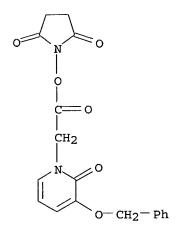
AB The title compds. [I (wherein X = S, O; Y = OR1, SR1; R1 = H, lower alkyl; W, V = OH, NO2, CF3, etc.), II (X = S, O; Y = OR1, SR1; R, R1 = H, lower alkyl, (CH2)nSH, etc.; n = 1-4), III (P, Q = H, lower alkyl, NH2CH2, etc.)], useful for therapy and prophylaxis of neurodegenerative disease such as Alzheimer's disease, were prepared Thus, treatment of 3-hydroxy-2(1H)-pyridinone with MeI in a sealed glass tube at 140° afforded 59% II [X = O; Y = OH; R = Me] which showed IC50 of 46 μM against tyrosine nitration and IC50 of 420 μM against lipid peroxidn.
IT 95215-73-9P

95215-73-9P
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
(Reactant or reagent)

(preparation of therapeutic antioxidants for Alzheimer's disease)

RN 95215-73-9 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[[[2-oxo-3-(phenylmethoxy)-1(2H)-pyridinyl]acetyl]oxy]- (9CI) (CA INDEX NAME)



L8 ANSWER 15 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1998:319625 HCAPLUS

DOCUMENT NUMBER: 129:54280

TITLE: Synthetic studies into 3-hydroxy-2(1)-pyridinone based

hexadentate metal(III) ion chelators

AUTHOR(S): Fox, Raymond C.; Taylor, Paul D.

CORPORATE SOURCE: Chemistry and Life Sciences, Research Triangle

Institute, Research Triangle Park, NC, 27709, USA Synthetic Communications (1998), 28(9), 1563-1574

SOURCE: Synthetic Communications (1998)
CODEN: SYNCAV; ISSN: 0039-7911

PUBLISHER: Marcel Dekker, Inc.

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 129:54280

AB An improved synthesis and purification of the hexadentate chelators, N,N,N,-tris[2-(3-hydroxy-2-oxo-1,2-dihydropyridin-1-

yl)acetamido]ethylamine and N,N,N,-tris[2-(3-hydroxy-4-methyl-2-oxo-1,2-dihydropyridin-1-yl)acetamido]ethylamine is described.

IT 208592-19-2P 208592-20-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of hydroxypyridinone-based hexadentate metal ion chelators)

RN 208592-19-2 HCAPLUS

CN 1H-Isoindole-1,3(2H)-dione, 2-[[(3-hydroxy-2-oxo-1(2H)-

pyridinyl)acetyl]oxy]- (9CI) (CA INDEX NAME)

RN 208592-20-5 HCAPLUS

CN 1H-Isoindole-1,3(2H)-dione, 2-[[(3-hydroxy-4-methyl-2-oxo-1(2H)-pyridinyl)acetyl]oxy]- (9CI) (CA INDEX NAME)

REFERENCE COUNT: 17 THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L8 ANSWER 16 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1997:805833 HCAPLUS

DOCUMENT NUMBER: 128:58278

TITLE: Nucleic acid hybridization using probes labeled with a

reporter group with spectroscopic properties sensitive

to hybrid formation

INVENTOR(S): Kubista, Mikael; Svanvik, Nicke

PATENT ASSIGNEE(S): Kubista, Mikael, Swed.; Svanvik, Nicke

SOURCE: PCT Int. Appl., 57 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PAT | ENT | NO. | | | KIN | D | DATE | | | APPL: | ICAT | ION 1 | NO. | | D | ATE | |
|-----|------|-----|-----|-----|-----|-----|------|----------|-----|-------|-------|-----------|-----|-----|-----|-----------|-----|
| WO | 9745 | 539 | | | A1 | _ | 1997 | 1204 | 1 | WO 1: | 997-: | SE95: | 3 | | 1: | 9970. | 530 |
| | W: | AL, | AM, | ΑT, | ΑU, | ΑZ, | BA, | BB, | BG, | BR, | BY, | CA, | CH, | CN, | CU, | CZ, | DE, |
| | | DK. | EE. | ES. | FI. | GB. | GE. | GH. | HU. | TL. | TS. | JP. | KE. | KG. | KP. | KR. | KZ. |

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LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL,
          PT, RO, RU, SD, SE, SG, SI, SK, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM

RW: GH, KE, LS, MW, SD, SZ, UG, AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN,
               ML, MR, NE, SN, TD, TG
                                     19971201
                                                   SE 1996-2183
     SE 9602183
                              Α
                                                                              19960531
     SE 506700
                              C2
                                     19980202
     CA 2256545
                              AA
                                     19971204
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     CA 2451442
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                                     19971204
                                                   CA 1997-2451442
                                                                              19970530
                                                   AU 1997-31129
     AU 9731129
                              A1
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                                                                              19970530
                                                   EP 1997-926344
                                                                              19970530
     EP 918852
                              A1
                                     19990602
          R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, NL, SE, PT, IE, FI
     BR 9709495
                              Α
                                     19990810
                                                   BR 1997-9495
                                                                              19970530
     CN 1226928
                              Α
                                     19990825
                                                   CN 1997-196872
                                                                              19970530
                                                   JP 1997-542246
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                              T2
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                              A1
                                                   EP 2003-11589
     EP 1357185
                                     20031029
                                                                              19970530
          R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, NL, SE, PT, IE, FI
     RU 2223966
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                                     20040220
                                                   RU 1998-123553
                                                                              19970530
     PL 188875
                              B1
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                                                   PL 1997-366875
                                                                              19970530
     US 6329144
                             B1
                                     20011211
                                                   US 1999-194679
                                                                              19990129
     AU 768137
                             B2
                                     20031204
                                                   AU 2001-55970
                                                                              20010725
                             A5
     AU 2001055970
                                     20010927
     JP 2005047901
                             A2
                                     20050224
                                                   JP 2004-204075
                                                                              20040712
PRIORITY APPLN. INFO.:
                                                   SE 1996-2183
                                                                          A 19960531
                                                   AU 1997-31129
                                                                          A3 19970530
                                                   CA 1997-2256545
                                                                          A3 19970530
                                                   JP 1997-542246
                                                                          A3 19970530
                                                   WO 1997-SE953
                                                                          W 19970530
                                                   EP 1997-926344
                                                                          A3 19971204
```

OTHER SOURCE(S): MARPAT 128:58278

AB Nucleic acid hybridization using probes labeled with a reporter group that changes its spectroscopic properties upon formation of a hybrid is described. The preferred reporter is an asym. cyanine dye. Use of such a probe allows the immediate or real time quantification of hybrid formation. The method can be used with nucleic acid probes or analogs such as peptide nucleic acids. A pair of suitable benzothiazol quinoline dyes were synthesized by standard chem and analogs with spacer arms prepared for

conjugation to nucleic acids. A conjugate of one of these dyes bound to an immobilized peptide nucleic acid probe showed a 45-fold increase in fluorescence upon formation of a hybrid. Fluorescence showed up to a 50-fold increase in quantum yield (range 8-50-fold) depending upon the probe used.

IT 200262-06-2P

RL: ARU (Analytical role, unclassified); PRP (Properties); SPN (Synthetic preparation); ANST (Analytical study); PREP (Preparation)

(as reporter group in nucleic acid hybridization; nucleic acid

hybridization using probes labeled with reporter group with spectroscopic properties sensitive to hybrid formation)

RN 200262-06-2 HCAPLUS

CN Benzothiazolium, 2-[[1-[2-[(2,5-dioxo-1-pyrrolidinyl)oxy]-2-oxoethyl]-4(1H)-pyridinylidene]methyl]-3-methyl- (9CI) (CA INDEX NAME)

L8 ANSWER 17 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1995:427460 HCAPLUS

DOCUMENT NUMBER: 123:83982

TITLE: Structure of cyclic hexa-pseudopeptide constructed

from N, N'-ethylene-bridged-(S)-alanyl-(S)-alanine and

glycine

AUTHOR(S): Kojima, Yoshitane; Yamashita, Tetsushi; Miyake,

Hiroyuki

CORPORATE SOURCE: Fac. Sci., Osaka City Univ., Osaka, 558, Japan

SOURCE: Chemistry Letters (1995), (3), 201-2

CODEN: CMLTAG; ISSN: 0366-7022

PUBLISHER: Nippon Kagakkai

DOCUMENT TYPE: Journal LANGUAGE: English

GI

I

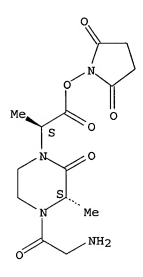
AB The crystal structure of 18-membered cyclic pseudopeptide I, containing N,N'-ethylene-bridged-(S)-alanyl-(S)-alanine and glycine was determined by x-ray crystallog. Moreover, the structure of this pseudopeptide was examined by 1H NMR measurement in CD3CN, and by mol. mechanics calcns.

IT 164857-03-8
 RL: RCT (Reactant); RACT (Reactant or reagent)
 (structure of cyclic hexapseudopeptide constructed from ethylene-bridged alanylalanine and qlycine)

RN 164857-03-8 HCAPLUS

CN Piperazinone, 4-(aminoacetyl)-1-[2-[(2,5-dioxo-1-pyrrolidinyl)oxy]-1-methyl-2-oxoethyl]-3-methyl-, monohydrochloride, [S-(R*,R*)]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



● HCl

L8 ANSWER 18 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1992:566123 HCAPLUS

DOCUMENT NUMBER: 117:166123

TITLE: Effect of the chemical modification by viologen on the

reduction of metmyoglobin

AUTHOR(S): Tsukahara, Keiichi; Todorobaru, Hiromi

CORPORATE SOURCE: Fac. Sci., Nara Women's Univ., Nara, 630, Japan

SOURCE: Chemistry Letters (1992), (7), 1181-4

CODEN: CMLTAG; ISSN: 0366-7022

DOCUMENT TYPE: Journal

LANGUAGE: English

AB Metmyoglobin covalently linked with viologen was prepared and reduced by dithionite ions faster than the native metmyoglobin, suggesting that the reduction by dithionite of the attached viologen was followed by a rapid intramol. electron transfer from the viologen radical cation to the heme iron center.

IT 143674-76-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and coupling of, with metmyoglobin)

RN 143674-76-4 HCAPLUS

CN 4,4'-Bipyridinium, 1-[2-[(2,5-dioxo-1-pyrrolidinyl)oxy]-2-oxoethyl]-1'-methyl-, diperchlorate (9CI) (CA INDEX NAME)

CM 1

CRN 143674-75-3 CMF C17 H17 N3 O4

CM 2

CRN 14797-73-0 CMF Cl O4

L8 ANSWER 19 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1990:229683 HCAPLUS

DOCUMENT NUMBER: 112:229683

TITLE: Novel 3-hydroxy-2(1H)-pyridinones. Synthesis,

iron(III)-chelating properties and biological activity
AUTHOR(S): Streater, Michael; Taylor, Paul D.; Hider, Robert C.;

Porter, John

CORPORATE SOURCE: Dep. Chem. Biol. Chem., Univ. Essex, Colchester, CO4

3SQ, UK

SOURCE: Journal of Medicinal Chemistry (1990), 33(6), 1749-55

CODEN: JMCMAR; ISSN: 0022-2623

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 112:229683

GI

AB The synthesis of a range of novel bidentate, e.g., I (R = alkyl or alkylaminocarbonylmethyl), and hexadentate ligands containing the chelating moiety 3-hydroxy-2(1H)-pyridinone is described. The pKa values of the ligands and the stability consts. of their iron(III) complexes were determined The stability constant of the iron(III) complex of one of the hexadentate ligands is comparable to that of desferrioxamine B. The distribution coeffs. of the ligands and their iron(III) complexes were also determined One of the novel hexadentate compds. markedly enhanced iron(III) excretion from both hepatocytes and iron-overloaded mice.

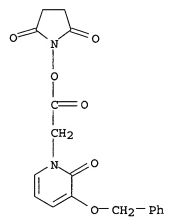
IT 95215-73-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reaction of, with amines)

RN 95215-73-9 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[[[2-oxo-3-(phenylmethoxy)-1(2H)-pyridinyl]acetyl]oxy]- (9CI) (CA INDEX NAME)



L8 ANSWER 20 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1990:7937 HCAPLUS

DOCUMENT NUMBER: 112:7937

TITLE: Preparation and testing of tripeptide derivatives as

cardiovascular agents

INVENTOR(S): Sawayama, Tadahiro; Nishimura, Kazuya; Deguchi,

Takashi

PATENT ASSIGNEE(S): Dainippon Pharmaceutical Co., Ltd., Japan

SOURCE: Jpn. Kokai Tokkyo Koho, 10 pp.

CODEN: JKXXAF

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|---------------------------------------|--------|----------|----------------------------------|----------------------|
| | | | | |
| JP 01125357 PRIORITY APPLN. INFO.: | A2 | 19890517 | JP 1987-281873 JP 1987-281873 | 19871106 19871106 |
| OTHER SOURCE(S): | MARPAT | 112:7937 | | |

$$Q = -(CH_2) mN$$

$$Q^1 = - CO_2 R^2$$

$$Q^2 = - CO_2 R^2$$

$$Q^3 = - CO_2 R^2$$

RR1CHCONHCH(CO2R2)(CH2)2COR3 [I; R = H, lower alkyl, PhCH2; R1 = AB (NH)m(CH2)nW, Q; R2 = H, lower alkyl; R3 = Q1, Q2, Q3, NR4CHR2CO2R2; W = H, CO2H, NH2, OH; Y = H, lower alkyl, Ph, PhCH2; R4 = C4-8 cycloalkyl, halo, alkoxy, (OH-substituted) Ph; m = 0, 1; n = 0-4] and their salts are prepared Refluxing 28 g 2-(S)-bromopropionic acid with 42 g PhCH2OH in PhMe gave 17.0 g benzyl 2-(S)-bromopropionate, 2.2 g of which was stirred with 1.6 g 1-benzylpiperazine in MeCN, then hydrolyzed with aqueous NaOH to give 1.0 g 2-(R)-(4-benzylpiperazinyl) propionic acid (II). Then, 24.5 g N-benzyloxycarbonyl-O1-ethyl-D-glutamic acid was stirred with 17.5 g Et (2S, 3aS, 7aS)-octahydro-1H-indole-2-carboxylate-HCl in CH2Cl2, then reduced, and then hydrolyzed with aqueous NaOH to give 15.01 g (2S, 3aS, 7aS)-1-(γ-D-glutamyl)octahydro-1H-indole-2-carboxylic acid (III). Then, 0.8 g II was treated with 0.4 g N-hydroxysuccinimide in CHCl3 to give 2-(R)-(4-benzylpiperazinyl)propionic acid N-hydroxysuccinimide ester, which was treated with 1.0 g III in THF to give 0.8 g (2S, 3aS, 7aS) -1- [N-2(R) - (4-benzylpiperazinyl) propionyl] - γ -Dqlutamyl]octahydro-1H-indole-2-carboxylic acid, 0.4 g of which was refluxed with HCO2H in MeOH in the presence of Pd black for 4 h to give 0.2 g (2S, 3aS, 7aS)-1-[N-(2R)-piperazinylpropionyl)- γ -Dglutamyl]octahydro-1H-indole-2-carboxylic acid, which showed an IC50 of 2.1 + 10-7 M against angiotensin converting enzyme.

IT 124078-64-4P

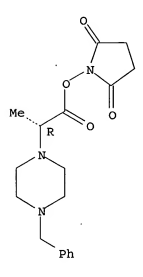
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and condensation of, with (glutamyl)indolecarboxylic acid)

RN 124078-64-4 HCAPLUS
CN 2,5-Pyrrolidinedione, 1-[1-oxo-2-[4-(phenylmethyl)-1-piperazinyl]propoxy]-

CN 2,5-Pyrrolidinedione, 1-[1-oxo-2-[4-(phenylmethyl)-1-piperazinyl]propoxy], (R)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.



L8 ANSWER 21 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1989:589581 HCAPLUS

DOCUMENT NUMBER:

111:189581

TITLE:

Morpholinoalkylcarboxylates as plant growth regulators

and fungicides

INVENTOR(S):

Ballschuh, Detlef; Banasiak, Lothar; Gruenzel, Hermann; Kluge, Eberhard; Lyr, Horst; Ohme, Roland;

Rusche, Jochen; Seibt, Horst; Spengler, Dieter;

Stoeckel, Christian

PATENT ASSIGNEE(S):

Akademie der Landwirtschaftwissenschaften der DDR, Institut fuer Pflanzenschutzforschung, Ger. Dem. Rep.

Ger. (East), 28 pp.

SOURCE: Ger. (East

CODEN: GEXXA8

DOCUMENT TYPE:

Patent German

LANGUAGE:

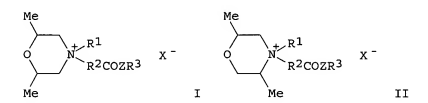
GΙ

Germ

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|--------|------------|-----------------|----------|
| | | | | |
| DD 263688 | A1 | 19890111 | DD 1985-278326 | 19850705 |
| PRIORITY APPLN. INFO.: | | | DD 1985-278326 | 19850705 |
| OTHER SOURCE(S): | MARPAT | 111:189581 | | |



AB Mixts. of the title compds. I and II [R1 = C6-20; R2 = C1-6 alkylene; R3 = (un)substituted alkyl, alkenyl, cycloalkyl, etc.; Z = 0, S; X- = anion]

(cis and/or trans) are prepared as fungicides and plant growth regulators. The fungicidal activity is both curative and preventive. Many target fungal species and host plants are listed. A mixture of cis- and/or trans-2,5-dimethyl-N-isotridecylmorpholine and cis- and/or trans-2,6-dimethyl-N-isotridecylmorpholine was refluxed with ClCH2CO2Me in NaI-containing acetonitrile, to give I-II (R1 = isotridecyl, R2 = CH2, R3 = Me, Z = 0, X = Cl).

IT 123322-75-8P 123322-78-1P 123340-64-7P 123340-67-0P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as fungicide and plant growth regulator)

RN 123322-75-8 HCAPLUS

CN Morpholinium, 4-isotridecyl-2,5-dimethyl-4-[2-(4-nitrophenoxy)-2-oxoethyl]-, chloride (9CI) (CA INDEX NAME)

● C1 -

RN 123322-78-1 HCAPLUS

CN Morpholinium, 4-[2-(2,6-dibromo-4-nitrophenoxy)-2-oxoethyl]-4-isotridecyl-2,5-dimethyl-, chloride (9CI) (CA INDEX NAME)

• cl -

RN 123340-64-7 HCAPLUS

CN Morpholinium, 4-isotridecyl-2,6-dimethyl-4-[2-(4-nitrophenoxy)-2-oxoethyl]-, chloride (9CI) (CA INDEX NAME)

• c1 -

RN 123340-67-0 HCAPLUS

CN Morpholinium, 4-[2-(2,6-dibromo-4-nitrophenoxy)-2-oxoethyl]-4-isotridecyl-2,6-dimethyl-, chloride (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{O}_2\text{N} & \text{Br} & \text{O} & \text{(C}_{13}\text{H}_27\text{-iso} \\ \text{O} & \text{C} & \text{CH}_2 & + \text{N} & \text{O} \\ \text{Br} & & \text{Me} \end{array}$$

Ocl-

L8 ANSWER 22 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1988:423356 HCAPLUS

DOCUMENT NUMBER:

109:23356

TITLE:

Interactions of organic substrates with 30- and

36-membered ring peptides containing

(2S,3'S)-2-(2'-oxo-3'-methylpiperazin-1'-yl)propanoic

acid and sarcosine

AUTHOR (S):

Kojima, Yoshitane; Yamashita, Tetsushi; Shibata, Kozo;

Ohsuka, Akio

CORPORATE SOURCE:

Fac. Sci., Osaka City Univ., Osaka, 558, Japan

SOURCE:

Polymer Journal (Tokyo, Japan) (1987), 19(10), 1221-3

CODEN: POLJB8; ISSN: 0032-3896

DOCUMENT TYPE:

Journal

LANGUAGE:

English

GI

AB Synthetic routes to cyclic peptides cyclo(Sar-EAA)4 (EAA = residue of title acid I) and cyclo(Sar-Sar-EAA)2 are described. Interaction of these cyclic peptides with p-toluenesulfonic acid salt of sodium, benzylamine, and 4-phenylbutylamine were studied by 1H NMR.

IT 114967-10-1P

CN

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and cyclization of)

RN 114967-10-1 HCAPLUS

1-Piperazineacetamide, N-[2-[4-[2-[[2-[4-[2-[(2,5-dioxo-1-pyrrolidinyl)oxy]-1-methyl-2-oxoethyl]-2-methyl-3-oxo-1-piperazinyl]-2-oxoethyl]methylamino]-1-methyl-2-oxoethyl]-2-methyl-3-oxo-1-piperazinyl]-2-oxoethyl]-N,α,3-trimethyl-4-[[methyl[2-[3-methyl-4-[(methylamino)acetyl]-2-oxo-1-piperazinyl]-1-oxopropyl]amino]acetyl]-2-oxo-, [3S-[1[R*[R*[R*[R*(R*)]]]],3R*,4[R*(R*)]]]-, mono(trifluoroacetate) (9CI) (CA INDEX NAME)

CM 1

CRN 114967-09-8 CMF C48 H73 N13 O15

PAGE 1-A

PAGE 1-B

CM 2

CRN 76-05-1 CMF C2 H F3 O2

ANSWER 23 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

1988:49290 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 108:49290

In vivo evaluation of hydroxypyridone iron chelators TITLE:

in a mouse model

AUTHOR (S): Gyparaki, M.; Porter, J. B.; Streater, M.; Hider, R.

C.; Huehns, E. R.

Dep. Haematol., Univ. Coll. London, London, WC1 E6HX, CORPORATE SOURCE:

UK

SOURCE: Acta Haematologica (1987), 78(3), 217-21

CODEN: ACHAAH; ISSN: 0001-5792

DOCUMENT TYPE:

Journal LANGUAGE: English

GI

Six N-substituted 3-hydroxypyrid-4-ones (I; R = Me, Et, Pr, iso-Pr, Bu, or AB hexyl) caused excretion when given i.p. to Fe-overloaded mice. The 1st 5 I were also active when given orally. Based on considerations of toxicity and relative activity, the compds. most promising for clin. use appeared to be I (R = Et) and I (R = Pr). A hexadentate pyrid-2-one (II) also caused Fe excretion when given i.p. or orally.

95215-73-9 IT

RL: RCT (Reactant); RACT (Reactant or reagent) (reaction of, with tris(aminoethyl)amine)

RN95215-73-9 HCAPLUS

2,5-Pyrrolidinedione, 1-[[[2-oxo-3-(phenylmethoxy)-1(2H)-CN

pyridinyl]acetyl]oxy] - (9CI) (CA INDEX NAME)

L8 ANSWER 24 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1987:156487 HCAPLUS

DOCUMENT NUMBER: 106:156487

TITLE: Salts of morpholinocarboxylic esters and

morpholinoalkyl phenyl ethers, processes for their preparation, and their use as fungicides and plant

growth regulators.

INVENTOR(S): Banasiak, Lothar; Leuner, Brita; Lyr, Horst; Nega,

Eva; Sunkel, Marianne

PATENT ASSIGNEE(S): Institut fuer Pflanzenschutzforschung Kleinmachnow,

Ger. Dem. Rep.

SOURCE: Eur. Pat. Appl., 41 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PAC | TENT NO. | | KIND | DATE | APPLICATION NO. | | DATE |
|----------|----------|---------|------------|-------------|-----------------|---|----------|
| | | | - - | | | | |
| EP | 209763 | | Al | 19870128 | EP 1986-108916 | | 19860701 |
| | R: AT, | BE, CH, | DE, | FR, GB, IT, | LI, LU, NL, SE | | |
| DD | 263685 | | A1 | 19890111 | DD 1985-278323 | | 19850705 |
| DD | 263687 | | A1 | 19890111 | DD 1985-278325 | | 19850705 |
| AU | 8659401 | | A1 | 19870108 | AU 1986-59401 | | 19860630 |
| DK | 8603151 | | Α | 19870106 | DK 1986-3151 | | 19860702 |
| FI | 8602851 | | Α | 19870106 | FI 1986-2851 | | 19860704 |
| ZA | 8605002 | | Α | 19870325 | ZA 1986-5002 | | 19860704 |
| JP | 62084065 | | A2 | 19870417 | JP 1986-156349 | | 19860704 |
| HU | 42288 | | A2 | 19870728 | HU 1986-2826 | | 19860704 |
| HU | 42286 | | A2 | 19870728 | HU 1986-2827 | | 19860704 |
| ES | 2001853 | | A6 | 19880701 | ES 1986-125 | | 19860704 |
| PL | 146362 | | B1 | 19890131 | PL 1986-260474 | | 19860704 |
| CS | 264279 | | B2 | 19890613 | CS 1986-5135 | | 19860707 |
| PRIORITY | APPLN. | INFO.: | | | DD 1985-278323 | Α | 19850705 |
| | | | | | DD 1985-278325 | Α | 19850705 |
| | | | | | | | |

GΙ

Me
$$R^1$$
 X^- Me I

AB The title compds. [I; R = C6-20 alkyl; R2 = R3Z1CO, (un)substituted PhO; R3 = (halo)alkenyl, alkynyl, (un)substituted alkyl, cycloalkyl, aryl, aralkyl; X1 = anion of a nonphytotoxic acid; Z = O, S; Z1 = C1-6 alkylene; R3 and X- may be absent] were prepared as fungicides and plant growth regulators. A mixture of 30 g 4-isotridecyl-2,6-dimethylmorpholine and 10.9 g ClCH2CO2Me was refluxed 20 h in MeCN containing catalytic NaI to give 38 g I (R1 = isotridecyl, R2 = CO2Me, X = C1, Z = CH2)(II). At 10 μg/mL II gave 88% inhibition of growth of Botrytis cinerea. At 1000 mg/L II reduced the growth of cucumber plants by 32%.

IT 107562-00-5DP, quaternary derivs. 107562-11-8DP,

quaternary derivs.

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as fungicide and plant growth inhibitor)

RN 107562-00-5 HCAPLUS

CN 4-Morpholineacetic acid, 2,6-dimethyl-, 4-nitrophenyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} O_2N & O & Me \\ \hline \\ O-C-CH_2-N & O \\ \hline \\ Me \end{array}$$

RN 107562-11-8 HCAPLUS

CN 4-Morpholineacetic acid, 2,6-dimethyl-, 2,6-dibromo-4-nitrophenyl ester (9CI) (CA INDEX NAME)

L8 ANSWER 25 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1985:596001 HCAPLUS

DOCUMENT NUMBER: 103:196001

TITLE: Hydroxypyridinone derivatives and pharmaceutical

compositions containing them

INVENTOR(S): Hider, Robert Charles; Kontoghiorghes, George; Silver,

Jack; Stockham, Michael Arthur

PATENT ASSIGNEE(S): National Research Development Corp., UK

SOURCE: Eur. Pat. Appl., 55 pp.

CODEN: EPXXDW

DOCUMENT TYPE:

LANGUAGE:

Patent English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| P | ATENT | NO. | | | KINI |) | DATE | | API | PLICATI | ои ио | | DATE |
|----------|--------|------|-----|-----|-----------|-----|-------|------|--------|---------|---------------|-------|----------|
| - | | | | | | - | | | | | | - | |
| | P 1384 | | | | | | | | EP | 1984-3 | 06438 | | 19840920 |
| _ | P 1384 | | | | A3 | | 19870 | | | | | | |
| E | P 1384 | 21 | | | B1 | | 19901 | L205 | | | | | |
| | R: | BE, | CH, | DE, | FR, | GB, | | - | NL, SI | | | | |
| U | S 4666 | 927 | | | Α | | 19870 | 519 | US | 1984-6 | 51684 | | 19840918 |
| G | B 2146 | 989 | | | A1 | | 19850 | 501 | GB | 1984-2 | 23799 | | 19840920 |
| G | B 2146 | 989 | | | B2 | | 19870 | 218 | | | | | |
| Z | A 8407 | 408 | | | Α | | 19860 | 528 | ZA | 1984-7 | 7408 | | 19840920 |
| E | P 3571 | 50 | | | A1 | | 19900 | 307 | EP | 1989-2 | 202213 | | 19840920 |
| E | P 3571 | 50 | | | B1 | | 19931 | L208 | | | | | |
| | R: | BE, | CH, | DE, | FR, | GB | , IT, | LI, | NL, SI | Ξ | | | |
| D | K 8404 | 536 | | | A | | 19850 | 324 | DK | 1984-4 | 1536 | | 19840921 |
| D | K 1583 | 49 | | | В | | 19900 | 507 | | | | | |
| | K 1583 | | | | | | 19901 | 1015 | | | | | |
| J | P 6009 | 4965 | | | A2 | | 19850 | 0528 | JP | 1984-2 | 201408 | | 19840925 |
| J | P 0600 | 2739 | | | B4 | | 19940 | 112 | | | | | |
| Ū | S 4863 | 913 | | | Α | | 19890 | 905 | US | 1986-9 | 944355 | | 19861222 |
| | S 4912 | | | | | | 19900 | 327 | US | 1986-9 | 944872 | | 19861222 |
| | S 5104 | | | | | | 19920 | 0414 | US | 1989-4 | 03054 | | 19890901 |
| | P 0219 | | | | | | 19900 | 727 | JP | 1989-3 | 324877 | | 19891212 |
| | P 0600 | | | | B4 | | 19940 | | | | | | |
| PRIORI | | | | . : | | | | | GB | 1983-2 | 25494 | Α | 19830923 |
| 11110111 | | | | • • | | | | | | | | | 19840918 |
| | | | | | | | | | | | | | 19840920 |
| | | | | | | | | | | | | | 19861222 |
| | | | | | | | | | 0.5 | 1700 2 | , , , , , , , | | 17001222 |

OTHER SOURCE(S):

CASREACT 103:196001

GI

$$^{R^{10}}$$
 O $^{NCH_{2}COR}$ II HO O $^{NCH_{2}CONHCH_{2}CH_{2}}$ N 3 III

AB Title compds. (I) were prepared, in which two or more 3-hydroxypyrid-2-one, 3-hydroxypyrid-4-one, or 1-hydroxypyrid-2-one rings are linked. Thus, 2,3-dihydroxypyridine and EtO2CCH2Br reacted to give hydroxypyridone II (R = OEt, R1 = H), which reacted with PhCH2Cl-NaOH to give 41% II (R = OH, R1 = CH2Ph). Treatment of the latter compound with DCC and N-hydroxysuccinimide gave 80% 1-succinimido ester, which condensed with N(CH2CH2NH2)3 to give, after hydrogenolysis, 68% triamide III. III removed 40% of iron from 59Fe(III)-loaded human transferrin, vs. 27% for EDTA. Addnl., the accumulation of the Fe(III) complex of III in human

erythrocytes was >9-fold that of Fe(III) citrate.

IT 95215-73-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and amidation of)

RN 95215-73-9 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[[[2-oxo-3-(phenylmethoxy)-1(2H)-pyridinyl]acetyl]oxy]- (9CI) (CA INDEX NAME)

L8 ANSWER 26 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1985:487778 HCAPLUS

DOCUMENT NUMBER:

103:87778

TITLE:

N-(1,2-Diacyl-2-halo-1-vinyl)pyridinium salts

INVENTOR(S):

Richter, Andreas M.; Fanghaenel, Egon

PATENT ASSIGNEE(S):

Technische Hochschule "Carl Schorlemmer"

Leuna-Merseburg, Ger. Dem. Rep.

SOURCE:

Ger. (East), 7 pp.

CODEN: GEXXA8

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE | |
|------------------------|------|----------|-----------------|----------|--|
| | | | | | |
| DD 215308 | A1 | 19841107 | DD 1983-251654 | 19830602 | |
| PRIORITY APPLN. INFO.: | | | DD 1983-251654 | 19830602 | |
| GI | | | | | |

$$\begin{array}{c|c}
R & COR^2 \\
NC = CR^3 & X^{-1}
\end{array}$$
R1CO I

AB Title compds. I (R = H, alkyl, alkoxy, dialkylamino; R1, R2 = alkyl, aralkyl, aryl, alkoxy, aralkoxy, aryloxy, alkylthio, aralkylthio, arylthio; R3 = F, Cl, Br, iodo; X- = halide, ClO4-, BF4-) were prepared by

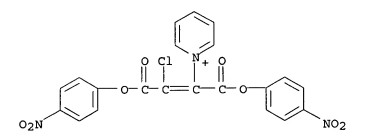
reacting R1COC.tplbond.CCOR2 with halogen and (un) substituted pyridines (II); or by reacting R1COCR4:CR5COR2 (R4, R5 = F, Cl, Br, iodo) with II; or by reacting R6COCR7:CR8COR9 (R6-R9 = F, Cl, Br, iodo) with R1OH or R1SH and II. Thus, 4-02NC6H4O2CCCl:CClCO2C6H4NO2-4 was treated with pyridine to give 99% I (R = H, R1 = R2 = 4-02NC6H4O, R3 = C1, X = C1). I are useful as intermediates in the preparation of dyes, heterocycles, polymers, and biol. active substances.

IT 97683-50-6P

> RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN

97683-50-6 HCAPLUS
Pyridinium, 1-[2-chloro-3-(4-nitrophenoxy)-1-[(4-nitrophenoxy)carbonyl]-3-CN oxo-1-propenyl]-, chloride (9CI) (CA INDEX NAME)



● cl -

ANSWER 27 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

1985:113302 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 102:113302

TITLE: Iron complexes for pharmaceutical compositions

INVENTOR(S): Hider, Robert Charles; Kontoghiorghes, George; Silver,

Jack; Stockham, Michael Arthur

PATENT ASSIGNEE(S): National Research Development Corp., UK

Brit. UK Pat. Appl., 19 pp. SOURCE:

CODEN: BAXXDU

DOCUMENT TYPE: Patent English LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|-----------|--------------|-----------------|------------|
| | | | | |
| GB 2136806 | A1 | 19840926 | GB 1984-7180 | 19840320 |
| GB 2136806 | B2 | 19870415 | | |
| EP 120670 | A1 | 19841003 | EP 1984-301882 | 19840320 |
| EP 120670 | B1 | 19881221 | | |
| R: BE, CH, DE, | FR, GE | , IT, LI, NL | , SE | |
| US 4650793 | Α | 19870317 | US 1984-592543 | 19840322 |
| DK 8401659 | Α | 19840925 | DK 1984-1659 | 19840323 |
| DK 159305 | В | 19901001 | | |
| DK 159305 | C | 19910304 | | |
| JP 59181258 | A2 | 19841015 | JP 1984-57186 | 19840324 |
| JP 06025120 | B4 | 19940406 | | |
| US 36831 | E | 20000822 | US 1995-390588 | 19950217 |
| PRIORITY APPLN. INFO.: | | | GB 1983-8055 | A 19830324 |

US 1984-592543

A5 19840322

OTHER SOURCE(S):

CASREACT 102:113302; MARPAT 102:113302

GΙ

OH OH OH II

AB Fe complexes of 3-hydroxy-2- and -4-pyridones I and II [R = acyl, (un)substituted hydrocarbon], useful for treatment of Fe deficiency anemia, were prepared Thus 2,3-dihydroxypyridine was treated with AcBr to give I (R = Ac), which complexed Fe(III) to give FeL3 (L = I; R = Ac) (III). Fe uptake from III by rat jejunal sacs in vitro was 38 times greater than from FeCl3.

IT 95215-73-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reactions of, with amines)

RN 95215-73-9 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[[[2-oxo-3-(phenylmethoxy)-1(2H)-pyridinyl]acetyl]oxy]- (9CI) (CA INDEX NAME)

L8 ANSWER 28 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1985:113301 HCAPLUS

DOCUMENT NUMBER:

102:113301

TITLE:

Pharmaceutical compositions

INVENTOR (S):

Hider, Robert Charles; Kontoghiorghes, George; Silver,

Jack

PATENT ASSIGNEE(S):

National Research Development Corp., UK

SOURCE:

Brit. UK Pat. Appl., 17 pp.

CODEN: BAXXDU

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PA | TENT NO. | | | KINI | | APPLICATION NO. | | DATE |
|---------|----------|------|-----|-----------|-------------|-----------------|----|----------|
| GB | 2136807 | | | A1 | 19840926 | GB 1984-7181 | | 19840320 |
| GB | 2136807 | | | | 19870423 | | | |
| CA | 1243606 | | | A1 | 19881025 | | | 19840207 |
| EP | 120669 | | | A2 | 19841003 | EP 1984-301881 | | 19840320 |
| EP | 120669 | | | A3 | 19850123 | | | |
| EP | 120669 | | | B1 | 19910123 | | | |
| | R: BE, | CH, | DE, | FR, | GB, IT, LI, | NL, SE | | |
| EP | 305646 | | | A2 | 19890308 | EP 1988-107000 | | 19840320 |
| EP | 305646 | | | A3 | 19900808 | | | |
| EP | 305646 | | | B1 | 19961113 | | | |
| | R: BE, | CH, | DE, | FR, | GB, IT, LI, | NL, SE | | |
| US | 4585780 | | | Α | | US 1984-592271 | | 19840322 |
| JP | 59205361 | | | A2 | 19841120 | JP 1984-57185 | | 19840324 |
| JP | 06072097 | | | B4 | 19940914 | | | |
| CA | 1338496 | | | A1 | 19960730 | CA 1986-524044 | | 19861128 |
| JP | 06080637 | | | A2 | 19940322 | JP 1993-150865 | | 19930622 |
| JP | 07064815 | | | B4 | 19950712 | | | |
| US | 35948 | | | E | 19981103 | US 1995-397321 | | 19950217 |
| PRIORIT | Y APPLN. | INFO | . : | | | GB 1983-8054 | Α | 19830324 |
| ·· | | | | | | CA 1984-446932 | A3 | 19840207 |
| | | | | | | EP 1984-301881 | P | 19840320 |
| | | | | | | US 1984-592271 | A5 | 19840322 |

GI

$$\bigcap_{R}^{OH} \bigcap_{I}^{O} \bigcap_{R}^{OH}$$

N-Substituted 3-hydroxy-2- or -4-pyridones I and II [R = acyl, (un)substituted hydrocarbon], useful for Fe complexation in vivo in treating Fe overloads, were prepared Thus 2,3-dihydroxypyridine was treated with AcBr to give hydroxy-2-pyridone I (R = Ac) (III). At 10 mg/mouse intragastrically, III gave 137 ± 45% excretion of 59Fe lactoferrin in Fe-loaded mice, compared to blank controls.

IT 95215-73-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reactions of, with amines)

RN 95215-73-9 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[[[2-oxo-3-(phenylmethoxy)-1(2H)-pyridinyl]acetyl]oxy]- (9CI) (CA INDEX NAME)

L8 ANSWER 29 OF 29 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1985:113157 HCAPLUS

DOCUMENT NUMBER: 102:113157

TITLE: Preparation and biological effects of

N-[tris(hydroxymethyl)methylaminocarbonylmethyl]

derivatives of heterocyclic bases

AUTHOR(S): Pischel, Helmut; Holy, Antonin; Vesely, Jiri; Wagner,

Guenther

CORPORATE SOURCE: Sekt. Biowiss.-Pharm., Karl-Marx-Univ., Leipzig, Ger.

Dem. Rep.

Journal

SOURCE: Collection of Czechoslovak Chemical Communications

(1984), 49(11), 2541-50

CODEN: CCCCAK; ISSN: 0366-547X

DOCUMENT TYPE:

LANGUAGE: English

GI

AB The title compds. were synthesized by the reaction of (HOCH2)3CNH2 with p-nitrophenyl or alkyl esters of N-carboxymethyl derivs. of uracil, 5-chloro-, 5-bromo-, 5-iodouracil, thymine, cytosine, 6-azauracil, 2-pyridone, 2-pyrimidone, 3-pyridazone and orotic acid. Some novel intermediates were also prepared Of all the amides tested, only the 3-pyridazone derivative I and orotic acid derivative II inhibited the growth of L-1210 mouse leukemic cells in vitro with ID50 .apprx.10-4 mol 1-1.

IT 95209-96-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

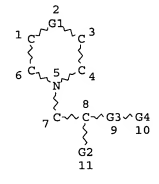
(preparation and reaction of, with TRIS)

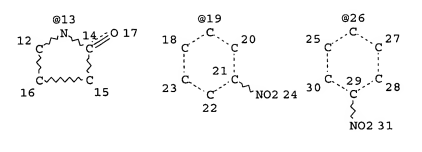
RN 95209-96-4 HCAPLUS

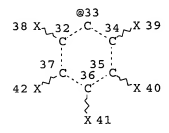
CN 1(2H)-Pyridineacetic acid, 2-oxo-, 4-nitrophenyl ester (9CI) (CA INDEX

NAME)

=> => d stat que .







VAR G1=C/N/O VAR G2=O/S/N VAR G3=O/S VAR G4=13/19/26/33 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 42

STEREO ATTRIBUTES: NONE

82 SEA FILE=REGISTRY SSS FUL L1 L5

L6 STR

NO2 31

VAR G1=C/N/O
VAR G2=O/S/N
VAR G3=O/S
VAR G4=13/19/26/33
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC 1

NUMBER OF NODES IS 42

STEREO ATTRIBUTES: NONE

L7 46 SEA FILE=REGISTRY SUB=L5 SSS FUL L6
L8 29 SEA FILE=HCAPLUS ABB=ON PLU=ON L7

L9 36 SEA FILE=REGISTRY ABB=ON PLU=ON L5 NOT L7

L10 18 SEA FILE=HCAPLUS ABB=ON PLU=ON L9

L11 17 SEA FILE=HCAPLUS ABB=ON PLU=ON L10 NOT L8

=> d ibib abs hitstr l11 1-17

L11 ANSWER 1 OF 17 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:399339 HCAPLUS

DOCUMENT NUMBER: 141:254556

TITLE: Grassland's locoweed toxin vaccine

INVENTOR(S): Dong, Dewen; Cao, Guangrong; Zhao, Baoyu; Ge, Pengbin PATENT ASSIGNEE(S): Danong Biotechnology Co., Ltd., Yangling, Peop. Rep.

Chi

SOURCE: Faming Zhuanli Shenging Gongkai Shuomingshu, 17 pp.

CODEN: CNXXEV

DOCUMENT TYPE: Patent LANGUAGE: Chinese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

CN 1395967 A 20030212 CN 2002-114592 20020524

PRIORITY APPLN. INFO.: CN 2002-114592 20020524

AB The process comprises N-alkylating swainsonine with bromoacetic acid N-succinimido ester in acetone under refluxing, coupling with bovine serum albumin in water at 0 °C, dialyzing, freeze drying, and emulsifying with Freund's adjuvant.

IT 754196-04-8P

RL: PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP
(Preparation); RACT (Reactant or reagent)
 (vaccine for Grassland's locoweed toxin)

RN 754196-04-8 HCAPLUS

CN Indolizinium, 4-[2-[(2,5-dioxo-1-pyrrolidinyl)oxy]-2-oxoethyl]octahydro-1,2,8-trihydroxy-, bromide, (1S,2R,8R,8aR)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

• Br-

L11 ANSWER 2 OF 17 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:913165 HCAPLUS

DOCUMENT NUMBER: 139:381472

TITLE: Preparation of naphthaldiimide derivatives as

anti-Helicobacter agents

INVENTOR(S): Sugimori, Giichi; Masui, Moriyasu; Nishida, Kuniyoshi;

Hasegawa, Yasushi; Kobayashi, Naotake

PATENT ASSIGNEE(S): Shionogi & Co., Ltd., Japan

SOURCE: PCT Int. Appl., 157 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|---------------|------|----------|-----------------|----------|
| | | | | |
| WO 2003095453 | A1 | 20031120 | WO 2003-JP5795 | 20030508 |

W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NI, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IT, LU, MC, NL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG **A1** 20031111 AU 2003-235908 AU 2003235908 20030508 PRIORITY APPLN. INFO.: JP 2002-137845 20020513 WO 2003-JP5795 W 20030508

OTHER SOURCE(S):

MARPAT 139:381472

Ι

GI

RN

AB Title compds. I (R1, R2 = H, alkyl, cycloalkyl, heterocyclyl, etc; X1, X2, Y1, Y2 = H, halo, etc.) are prepared When employed alone, such a compound is useful as an agent against Helicobacter. When employed as a combined drug, it can remarkably lessen side effects occurring in treating digestive ulcer, etc. These compds. or compns. can specifically injure and remove Helicobacter to thereby effectively treat digestive diseases (for example, gastric ulcer, duodenal ulcer, gastritis and gastric cancer).

IT 625085-55-4P 625086-14-8P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of naphthaldiimide derivs. as anti-Helicobacter agents) 625085-55-4 HCAPLUS

CN Benzo[lmn][3,8]phenanthroline-2(1H)-acetic acid, 3,6,7,8-tetrahydro-1,3,6,8-tetraoxo-7-(2-pyridinyl)-, 4-nitrophenyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ N & & & \\ \hline \end{array}$$

625086-14-8 HCAPLUS RN

Benzo[lmn][3,8]phenanthroline-2(1H)-acetic acid, 3,6,7,8-tetrahydro-CN 1,3,6,8-tetraoxo-7-(2-pyridinyl)-, 3-nitrophenyl ester (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & & \\ & & & \\ N & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ & & & \\ &$$

REFERENCE COUNT: 24 THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 3 OF 17 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2003:719471 HCAPLUS

DOCUMENT NUMBER:

139:261174

TITLE:

Preparation of N-heterocyclyl indole-2-carboxamides as

glycogen phosphorylase inhibitors

INVENTOR (S):

Birch, Alan Martin; Morley, Andrew David Astrazeneca AB, Swed.; Astrazeneca UK Limited

PATENT ASSIGNEE(S):

PCT Int. Appl., 86 pp.

SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO. | | DATE APPLICATION NO. | | | | |
|------------------------|-----------------|----------------------|-----------------|--|--|--|
| | | | | | | |
| WO 2003074513 | A2 20030912 | WO 2003-GB893 | 20030304 | | | |
| WO 2003074513 | A3 20031231 | | | | | |
| W: AE, AG, AL, | AM, AT, AU, AZ, | BA, BB, BG, BR, BY, | BZ, CA, CH, CN, | | | |
| CO, CR, CU, | CZ, DE, DK, DM, | DZ, EC, EE, ES, FI, | GB, GD, GE, GH, | | | |
| GM, HR, HU, | ID, IL, IN, IS, | JP, KE, KG, KP, KR, | KZ, LC, LK, LR, | | | |
| LS, LT, LU, | LV, MA, MD, MG, | MK, MN, MW, MX, MZ, | NO, NZ, OM, PH, | | | |
| PL, PT, RO, | RU, SC, SD, SE, | SG, SK, SL, TJ, TM, | TN, TR, TT, TZ, | | | |
| | UZ, VC, VN, YU, | | | | | |
| | | SL, SZ, TZ, UG, ZM, | ZW. AM. AZ. BY. | | | |
| | | BE, BG, CH, CY, CZ, | | | | |
| , , , | | LU, MC, NL, PT, RO, | | | | |
| , , , | • | GN, GQ, GW, ML, MR, | | | | |
| AU 2003216991 | | AU 2003-216991 | | | | |
| | | EP 2003-712313 | | | | |
| | | GB, GR, IT, LI, LU, | | | | |
| | | CY, AL, TR, BG, CZ, | | | | |
| • • • • | | US 2003-506748 | • | | | |
| | | | | | | |
| | 12 20050825 | JP 2003-572981 | | | | |
| PRIORITY APPLN. INFO.: | | GB 2002-5162 | | | | |
| | | WO 2003-GB893 | w 20030304 | | | |
| OTHER SOURCE(S): | MARPAT 139:2611 | 74 | | | | |
| GT | | | | | | |

GI

$$\begin{bmatrix} \mathbb{R}^4 \end{bmatrix}_{\mathfrak{m}} \xrightarrow{N}_{\mathfrak{m}} \begin{bmatrix} \mathbb{R}^2 \\ \mathbb{N} \end{bmatrix}_{\mathfrak{m}} \xrightarrow{\mathbb{R}^2}_{\mathfrak{m}} \begin{bmatrix} \mathbb{R}^1 \\ \mathbb{N} \end{bmatrix}_{\mathfrak{m}}$$

$$C1$$
 N
 N
 O
 O
 O
 O
 O
 O
 O

The title compds. [I; A = phenylene or heteroarylene; m = 0-2; n = 0-2; R1 = halo, NO2, CN, OH, CO2H, etc.; R2 = H, OH, CO2H; R3 = H, OH, aryl, heterocyclyl, etc.; R4 = H, halo, NO2, CN, etc.] which possess glycogen phosphorylase inhibitory activity and accordingly have value in the treatment of disease states associated with increased glycogen phosphorylase activity such as diabetes type II, were prepared Thus, amidation of 5-chloro-1H-indole-2-carboxylic acid with Me 2-(3-amino-2-oxo-3,4-dihydroquinolin-1-(2H)-yl)acetate (preparation given) in the presence of HOBT, DCM and EDCI afforded 59% II. The compds. I showed IC50 values in the range 100μM to 1nM against against hrl glycogen phosphorylase a. Pharmaceutical composition comprising the compound I was claimed.

IT 599193-13-2P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of N-heterocyclyl indole-2-carboxamides as glycogen phosphorylase inhibitors)

RN 599193-13-2 HCAPLUS

CN 1H-Indole-2-carboxamide, 5-chloro-N-[1-[2-[(2,5-dioxo-1-pyrrolidinyl)oxy]-2-oxoethyl]-1,2,3,4-tetrahydro-2-oxo-3-quinolinyl]- (9CI) (CA INDEX NAME)

L11 ANSWER 4 OF 17 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2002:575812 HCAPLUS

DOCUMENT NUMBER: 137:381573

TITLE: Chemical Ribonucleases: 4.1 An Analysis of the Domain

Structure of Chemical Ribonucleases Based on

1,4-Diazabicyclo[2.2.2]octane

AUTHOR(S): Konevetz, D. A.; Mironova, N. L.; Beck, I. E.;

Zenkova, M. A.; Shishkin, G. V.; Vlassov, V. V.;

Silnikov, V. N.

CORPORATE SOURCE: Novosibirsk Institute of Bioorganic Chemistry, Russian

Academy of Sciences, Siberian Branch, Novosibirsk,

630090, Russia

SOURCE: Russian Journal of Bioorganic Chemistry (Translation

of Bioorganicheskaya Khimiya) (2002), 28(4), 331-341

CODEN: RJBCET; ISSN: 1068-1620

PUBLISHER: MAIK Nauka/Interperiodica Publishing

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 137:381573

Artificial RNases of the ABLkCm series were synthesized. They consist of a lipophilic alkyl radical (Et, n-C14H29, or n-C15H31) [Acy], an "RNA-binding domain" [Vcy] (bisquaternary salt of 1,4diazabicyclo[2.2.2]octane), a "catalytic domain" [Scy]m [histamine ([Scy]1) or histidine ([Scy]3) residue], and a "linker" Lk that joins the "domains" B and Cm [here, k is the number of methylene units (one or three) in the linker]. The effect of the "domain structure" on the catalytic properties of the chemical RNases was analyzed using seven compds. of this series (ABL1C1, ABL3C1, ABL3C3, AC1, AB, BL2, and BL3C3). The catalytic activity of the compds. was assessed in the reaction of hydrolysis of the in vitro transcripts of human tRNALys and yeast tRNAAsp under physiol. conditions. It was shown that only chemical RNases that involve all the fragments of the ABLkCm construct can hydrolyze the substrate tRNA at a high rate (90% of tRNA is hydrolyzed for 10 h at 37°[Scy]). The activity of the compds. is largely determined by the presence of a long lipophilic radical linked to 1,4-diazabicyclo[2.2.2]octane and a long linker, which joins the RNA-hydrolyzing and RNA-binding domains. results indicate an important role of hydrophobic interactions in the acceleration of the RNA hydrolysis reaction.

IT 475661-85-9P

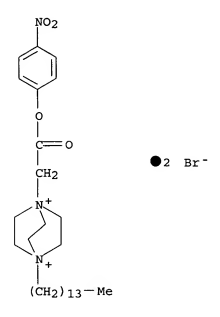
RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT

(Reactant or reagent)

(all domains of chemical RNase are required for efficient tRNA hydrolysis)

RN 475661-85-9 HCAPLUS

CN 1,4-Diazoniabicyclo[2.2.2]octane, 1-[2-(4-nitrophenoxy)-2-oxoethyl]-4-tetradecyl-, dibromide (9CI) (CA INDEX NAME)



REFERENCE COUNT: 24 THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

RECORD. AND CITATIONS AVAILABLE IN THE RE FORMAL

L11 ANSWER 5 OF 17 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2002:504749 HCAPLUS

DOCUMENT NUMBER: 137:79227

TITLE: Novel functional peptide nucleic acid monomer and

process for producing the same

INVENTOR(S): Ikeda, Hisafumi; Saito, Isao; Kitagawa, Fumihiko

PATENT ASSIGNEE(S): Applied Biosystems Japan Ltd., Japan

SOURCE: PCT Int. Appl., 63 pp.

CORCE: FCI IIIC. Appl., 03

CODEN: PIXXD2
OCUMENT TYPE: Patent

DOCUMENT TYPE: Patent LANGUAGE: Japanese

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

GI

| PATENT NO. | KIND DATE | E AP | PLICATION NO. | | DATE |
|------------------------|------------|-------------|-----------------|-------|------------|
| | | | | | |
| WO 2002051797 | A1 2002 | 20704 WO | 2001-JP8120 | | 20010919 |
| W: JP, US | | | | | |
| RW: AT, BE, CH, | CY, DE, DK | , ES, FI, F | R, GB, GR, IE, | IT, L | U, MC, NL, |
| PT, SE, TR | | | | | |
| EP 1357112 | A1 2003 | 31029 EP | 2001-970133 | | 20010919 |
| R: AT, BE, CH, | DE, DK, ES | , FR, GB, G | R, IT, LI, LU, | NL, S | E, MC, PT, |
| IE, FI, CY, | TR | | | | |
| US 2004101839 | A1 2004 | 40527 US | 2003-250592 | | 20031224 |
| PRIORITY APPLN. INFO.: | | JP | 2000-394669 | Α | 20001226 |
| | | WO | 2001-JP8120 | W | 20010919 |
| OTHER SOURCE(S): | CASREACT 1 | 37:79227; M | ARPAT 137:79227 | 7 | |

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

```
A peptide nucleic acid (PNA) monomer represented by the following general
    formula A-(CH2)nCO-B [I; wherein A = Q or Q1 (wherein X = OH, Z = O; X =
    NH2, Z = H2N+; or X = NMe2, Z = Me2N+), Q2, Q3, Q4 (wherein R = hydrogen,
    NO2, NH2, NHCbz, bromine, fluorine, chlorine, or SO3Na2), Q5,
    3-(4-dimethylaminophenylazo)phenyl, 4-(4-dimethylaminophenylazo)phenylsulf
    onylamino, 2-(4-hydroxyphenylazo)benzoylamino, 5-
    dimethylaminonaphthalenesulfonylamino, 1-pyrenecarbonyl, 1-pyrenylmethyl,
    1-pyrenesulfonylamino, 6,7,8-trimethyl-1,3-dioxo-2,5-dihydro-2,4-
    diazaphenazin-2-yl, 4-methylcoumarin-7-ylaminocarbonyl,
    4-trifluoromethylcoumarin-7-ylaminocarbonyl, 4-methyl-2-oxo-1,2-
    dihydroquinoin-7-ylaminocarbonyl, 2-oxo-1,2-dihydroquinoin-3-
    ylaminocarbonyl, etc.; B is OH, pentafluorophenyloxy, succinimidyloxy,
    N-carboxylmethyl-N-[2-(tert-butoxycarbonylamino)ethyl]amino; n = an
    integer of 1 to 4] is prepared A PNA monomer I [A, N = same as above; B =
    N-carboxylmethyl-N-[2-(tert-butoxycarbonylamino)ethyl]amino] is prepared by
    amidation of an active ester I (A, n = same as above; B =
    pentafluorophenyloxy, succinimidyloxy) with tert-
    butoxycarbonylaminoethylamine or an \omega-amino acid derivative, in
    particular 2-[N-[2-(tert-butoxycarbonylamino)ethyl]amino]acetic acid (II).
    This process is convenient for the preparation of a photofunctional PNA monomer
    which is unstable under alkali condition. Thus, to a solution of 100 mg
    2-(5,7,8-trimethyl-1,3-dioxo-2,5-dihydro-2,4-diazaphenazin-2-yl)acetic
    acid and 70.2 mg pentafluorophenol in 10 mL DMF was added 73.2 mg
    1-ethyl-3-(3-dimethylaminopropyl)carbodiimide hydrochloride (EDC) at
     0° and stirred at 0° for 1 h and at room temperature for 12 h to
    give 85% 2,3,4,5,6-pentafluorophenyl 2-(5,7,8-trimethyl-1,3-dioxo-2,5-
    dihydro-2,4-diazaphenazin-2-yl)acetate (III). To a solution of the active
    ester III (100 mg) and 45.4 mg II in 10 mL DMF was added 36.3 \mu L
    diisopropylethylamine and stirred at room temperature for 15 h to give 85%
     2-[N-[2-(tert-butoxycarbonylamino)ethyl]-2-[(5,7,8-trimethyl-1,3-dioxo-2,5-
    dihydro-2,4-diazaphenazin-2-yl)acetyl]amino]acetic acid.
IT
    439913-28-7P, [1,3-Dioxo-1H,3H-benz[de]isoquinolin-2-yl]acetic
    acid pentafluorophenyl ester 439913-30-1P, [5-Nitro-1,3-dioxo-
     1H, 3H-benz[de]isoquinolin-2-yl]acetic acid pentafluorophenyl ester
     439913-33-4P
    RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
     (Reactant or reagent)
        (preparation of novel functional peptide nucleic acid monomers by amidation
        of active esters with \alpha-[N-[\beta-(tert-
       butoxycarbonylamino)ethyl]amino]acetic acid.)
     439913-28-7 HCAPLUS
RN
     1H-Benz[de]isoquinoline-2(3H)-acetic acid, 1,3-dioxo-, pentafluorophenyl
CN
     ester (9CI) (CA INDEX NAME)
```

RN 439913-30-1 HCAPLUS

CN 1H-Benz[de]isoquinoline-2(3H)-acetic acid, 5-nitro-1,3-dioxo-, pentafluorophenyl ester (9CI) (CA INDEX NAME)

RN 439913-33-4 HCAPLUS

CN 1H-Benz[de]isoquinoline-2(3H)-acetic acid, 5-(acetylamino)-1,3-dioxo-, pentafluorophenyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT: 14 THERE ARE 14 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 6 OF 17 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2000:871429 HCAPLUS

DOCUMENT NUMBER: 134:189912

TITLE: Chemical ribonucleases: 3. The synthesis of organic

catalysts for the phosphodiester bond hydrolysis on

the basis of quaternary salts of 1,4-

diazabicyclo[2.2.2]octane

AUTHOR(S): Konevetz, D. A.; Beck, I. E.; Sil'nikov, V. N.;

Zenkova, M. A.; Shishkin, G. V.

CORPORATE SOURCE: Novosibirsk Institute of Bioorganic Chemistry,

Siberian Division, Russian Academy of Sciences,

Novosibirsk, 630090, Russia

SOURCE: Russian Journal of Bioorganic Chemistry (Translation

of Bioorganicheskaya Khimiya) (2000), 26(11), 765-773

CODEN: RJBCET; ISSN: 1068-1620

PUBLISHER: MAIK Nauka/Interperiodica

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 134:189912

On the basis of imidazole and bisquaternary salts of 1,4-diazabicyclo[2.2.2]octane, a number of highly effective catalysts of the nDm series (here, n is the number of pos. charges at neutral pH values and m is the digital code of the catalytically active fragment: 1, histamine, and 2, histidine Me ester) were synthesized for the cleavage of phosphodiester bonds in ribonucleic acids. A general method for the synthesis of chemical RNases was suggested, which helps vary both the number of pos. charges in their RNA-binding domain and the catalytic center. By the example of hydrolysis under physiol. conditions of the in vitro transcript of tRNALys from human mitochondria, it was shown that the RNA cleavage rate with the nDm conjugates increases approx. 30-fold along with the increase in the number of pos. charges from two to four.

IT 327189-89-9P 327189-91-3P 327189-96-8P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of artificial RNases for phosphodiester bond hydrolysis of RNA

on basis of quaternary salts of 1,4-diazabicyclo[2.2.2]octane)

RN 327189-89-9 HCAPLUS

CN 1,4-Diazoniabicyclo[2.2.2]octane, 1-[2-(1,3-dihydro-1,3-dioxo-2H-isoindol-2-yl)ethyl]-4-[2-(4-nitrophenoxy)-2-oxoethyl]-, dibromide (9CI) (CA INDEX NAME)

PAGE 1-A

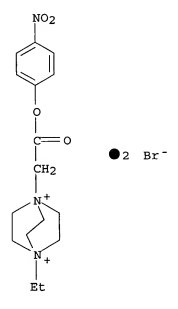
PAGE 2-A

●2 Br-

RN 327189-91-3 HCAPLUS

1,4-Diazoniabicyclo[2.2.2]octane, 1-(3-azidopropyl)-4-[2-(4-nitrophenoxy)-2-oxoethyl]-, bromide chloride (9CI) (CA INDEX NAME)

RN 327189-96-8 HCAPLUS
CN 1,4-Diazoniabicyclo[2.2.2]octane, 1-ethyl-4-[2-(4-nitrophenoxy)-2-oxoethyl]-, dibromide (9CI) (CA INDEX NAME)



REFERENCE COUNT: 19 THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 7 OF 17 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2000:735652 HCAPLUS

DOCUMENT NUMBER: 133:360397

TITLE: Chemical ribonucleases: 2. Design and hydrolytic activity of the ribonuclease mimics on the basis of

diazabicyclo[2.2.2] octane with a differing number of

positive charges

Zenkova, M. A.; Vlassov, A. V.; Konevets, D. A.; AUTHOR (S):

Silnikov, V. N.; Giege, R.; Vlassov, V. V.

Novosibirsk Institute of Bioorganic Chemistry, CORPORATE SOURCE:

Siberian Division, Russian Academy of Sciences,

Novosibirsk, 630090, Russia

Russian Journal of Bioorganic Chemistry (Translation SOURCE:

of Bioorganicheskaya Khimiya) (2000), 26(9), 610-615

CODEN: RJBCET; ISSN: 1068-1620

MAIK Nauka/Interperiodica

PUBLISHER: Journal DOCUMENT TYPE:

English LANGUAGE:

A procedure was proposed allowing one to synthesize RNase mimics on the basis of conjugates of diazabicyclo[2.2.2] octane with imidazole bearing a varying number of pos. charges (nDm series, where n is the number of pos. charges at neutral pH, m is the code of an imidazole-containing fragment of the catalytic domain: 1, histamine; 2, histidine Me ester). The hydrolytic activity of six compds. of this series was studied in physiol. conditions using in vitro transcript of human mitochondrial tRNALys as a substrate. It was shown that the rate of RNA hydrolysis with nDm conjugates rises with an increase in the number of pos. charges: an approx. 30-fold acceleration of hydrolysis was observed with an increase in the total charge of the construct from +2 to +4.

307305-05-1P 307305-06-2P IT

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(design and hydrolytic activity of RNase mimics based on diazabicyclo[2.2.2]octane and containing various number of pos. charges)

RN307305-05-1 HCAPLUS

1,4-Diazoniabicyclo[2.2.2]octane, 1-ethyl-4-[2-(4-nitrophenoxy)-2-CNoxoethyl] - (9CI) (CA INDEX NAME)

307305-06-2 HCAPLUS RN

1,4-Diazoniabicyclo[2.2.2]octane, 1-(3-azidopropyl)-4-[2-(4-nitrophenoxy)-CN2-oxoethyl]- (9CI) (CA INDEX NAME)

REFERENCE COUNT: 20 THERE ARE 20 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 8 OF 17 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2000:157026 HCAPLUS

DOCUMENT NUMBER: 133:4837

TITLE: Synthesis of a netropsin conjugate of a water-soluble

epi-quinocarcin analogue: the importance of

stereochemistry at nitrogen

AUTHOR(S): Herberich, B.; Scott, J. D.; Williams, R. M.

CORPORATE SOURCE: Department of Chemistry, Colorado State University,

Fort Collins, CO, USA

SOURCE: Bioorganic & Medicinal Chemistry (2000), 8(3), 523-532

CODEN: BMECEP; ISSN: 0968-0896

PUBLISHER: Elsevier Science Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

GI

AB The efficient synthesis of a water-soluble C11a-epi-analog I of quinocarcin is described. This substance, and a netropsin amide conjugate II lack the capacity to inflict oxidative damage on DNA due to the stereoelectronic geometry of their oxazolidine nitrogen atoms. The capacity of these substances to alkylate DNA through the generation of an iminium species has been examined Both compds. were found to be unreactive as DNA alkylating agents. The results of this study are discussed in the context of previous proposals on the mode of action of this family of antitumor alkaloids.

IT 165253-50-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(synthesis of a netropsin conjugate of a water-soluble epi-quinocarcin analog and the importance of stereochem. at nitrogen)

RN 165253-50-9 HCAPLUS

CN 2-Oxa-4,10c-diazaaceanthrylene-4(1H)-acetic acid, 2a,3,5,5a,6,10b-hexahydro-10-methoxy-3,3-dimethyl-, 4-nitrophenyl ester, (2aR,5aS,10bR)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

REFERENCE COUNT:

THERE ARE 35 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

II

L11 ANSWER 9 OF 17 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1999:759500 HCAPLUS

DOCUMENT NUMBER: 132:148595

Characterization and application of TITLE:

> acridine-9-N-acetyl-N-hydroxysuccinimide as a pre-column derivatization agent for fluorimetric detection of amino acids in liquid chromatography You, Jinmao; Lao, Wenjian; You, Jing; Wang, Guojun

Lanzhou Inst. Chem. Phys., Chinese Academy of Sciences, Lanchou, 730000, Peop. Rep. China CORPORATE SOURCE:

SOURCE:

Analyst (Cambridge, United Kingdom) (1999), 124(12),

1755-1760

CODEN: ANALAO; ISSN: 0003-2654 Royal Society of Chemistry

DOCUMENT TYPE: Journal English LANGUAGE:

AUTHOR(S):

PUBLISHER:

A simple and sensitive LC method that rapidly labels amino compds. including amino acids, using acridine-9-N-acetyl-N-hydroxysuccinimide (AAHS) which was synthesized by the reaction of acridine-9-N-acetic acid with benzenedisulfonyl-N-hydroxysuccinimide, was developed. A mixture of amines is treated with AAHS in the presence of triethylamine in non-aqueous acetonitrile or in 0.2 mol 1-1 borate buffer at pH 8.0-9.0 in 40% volume/volume acetonitrile solution to give quant. yields of amides. emission maximum for the derivatized amines is 435 nm ($\lambda ex = 404 \text{ nm}$). The labeled derivs. are very stable; no significant decomposition is observed after heating in 50% acetonitrile at 40° for 24 h. Studies on the derivatization conditions indicate that amines or amino acids react very rapidly with AAHS under the proposed conditions. The method, in conjunction with a multi-step gradient, offers baseline resolution of common amine or amino acid derivs. on a reversed-phase C18 column. This method is more convenient and more efficient than previous methods which require prior conversion of carboxylic acids to acyl chlorides, which are unstable to moisture. The LC separation of amine or amino acid derivs. has good reproducibility. The established method is also suitable for the determination of

other amine compds. in various biol. fluids.

150321-96-3P IT

RL: ARG (Analytical reagent use); SPN (Synthetic preparation); ANST (Analytical study); PREP (Preparation); USES (Uses)

(characterization and application of acridine-9-N-acetyl-Nhydroxysuccinimide as a pre-column derivatization agent for fluorimetric detection of amino acids in liquid chromatog.)

150321-96-3 HCAPLUS RN

2,5-Pyrrolidinedione, 1-[[(9-oxo-10(9H)-acridinyl)acetyl]oxy]- (9CI) CN INDEX NAME)

REFERENCE COUNT: 22 THERE ARE 22 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 10 OF 17 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1997:106528 HCAPLUS

DOCUMENT NUMBER: 126:212075

TITLE: Synthesis and chemiluminescent property of the novel

1,2-dioxetanes containing an acridane-10-acetate

moiety as the luminophor and trigger unit

AUTHOR(S): Imanishi, Takeshi; Ueda, Yohko; Tainaka, Ryoh;

Miyashita, Kazuyuki; Hoshino, Nobuhiro

CORPORATE SOURCE: Faculty Pharmaceutical Sciences, Osaka Univ., Suita,

EGE Tanan

565, Japan

SOURCE: Tetrahedron Letters (1997), 38(5), 841-844

CODEN: TELEAY; ISSN: 0040-4039

PUBLISHER: Elsevier
DOCUMENT TYPE: Journal

LANGUAGE: English

GI

I

AB Novel dioxetane derivs. I [R = Et, CH2CCl3, (un)substituted Ph] with an acridane-10-acetate moiety were prepared and tested as potential

chemiluminescent probes. The 10-acetate was found to play an important role both in stabilization and in base-mediated smooth degradation of the dioxetane ring.

IT 178312-97-5P 188002-48-4P

RL: PEP (Physical, engineering or chemical process); PRP (Properties); RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); PROC (Process); RACT (Reactant or reagent)

(preparation, thermal stability, and chemiluminescence of 1,2-dioxetanes containing an acridane acetate moiety)

RN 178312-97-5 HCAPLUS

CN Dispiro[acridine-9(10H),3'-[1,2]dioxetane-4',2''-

tricyclo[3.3.1.13,7]decane]-10-acetic acid, 4-nitrophenyl ester (9CI) (CA INDEX NAME)

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RN

CN Dispiro[acridine-9(10H),3'-[1,2]dioxetane-4',2''tricyclo[3.3.1.13,7]decane]-10-acetic acid, 2,4-dinitrophenyl ester (9CI)
(CA INDEX NAME)

PAGE 1-A

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IT 178313-00-3P 188002-39-3P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation, thermal stability, and chemiluminescence of 1,2-dioxetanes containing an acridane acetate moiety)

RN 178313-00-3 HCAPLUS

CN 10(9H)-Acridineacetic acid, 9-tricyclo[3.3.1.13,7]decylidene-, 4-nitrophenyl ester (9CI) (CA INDEX NAME)

RN 188002-39-3 HCAPLUS
CN 10(9H)-Acridineacetic acid, 9-tricyclo[3.3.1.13,7]decylidene-, 2,4-dinitrophenyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT: 27 THERE ARE 27 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L11 ANSWER 11 OF 17 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1996:401584 HCAPLUS

DOCUMENT NUMBER: 125:58346

Preparation of acridine derivatives as TITLE:

chemiluminescent compounds

INVENTOR (S): Imanishi, Takeshi; Hoshino, Nobuhiro; Shimamoto,

Kazutoshi

PATENT ASSIGNEE(S): Iatron Lab., Japan; Mitsubishi Chemical Yatron Co.,

Ltd.

SOURCE: Jpn. Kokai Tokkyo Koho, 10 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent

LANGUAGE:

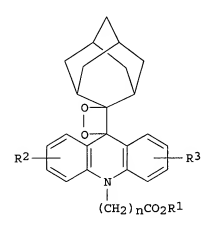
GΙ

Japanese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|--------|-----------|-----------------|----------|
| | | | | |
| JP 08092254 | A2 . | 19960409 | JP 1994-254730 | 19940923 |
| JP 3551984 | B2 | 20040811 | | |
| PRIORITY APPLN. INFO.: | | | JP 1994-254730 | 19940923 |
| OTHER SOURCE(S): | MARPAT | 125:58346 | | |



The title compds. I [n = 1 - 3; R1 = H, alkyl, etc.; R2, R3 = H, nitro, etc.] are prepared I [R2 = R3 = H; n = 1; R1 = 4-nitrophenyl] (II) (preparation

given) showed chemiluminescence. II showed good storage stability.

IT 178313-00-3P

> RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of acridine derivs. as chemiluminescent compds.)

RN178313-00-3 HCAPLUS

CN10(9H)-Acridineacetic acid, 9-tricyclo[3.3.1.13,7]decylidene-, 4-nitrophenyl ester (9CI) (CA INDEX NAME)

Ι

178312-96-4P 178312-97-5P ITRL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological

study); PREP (Preparation); USES (Uses)

(preparation of acridine derivs. as chemiluminescent compds.)

178312-96-4 HCAPLUS RN

CN

Dispiro[acridine-9(10H),3'-[1,2]dioxetane-4',2''-tricyclo[3.3.1.13,7]decane]-10-acetic acid, pentafluorophenyl ester (9CI) (CA INDEX NAME)

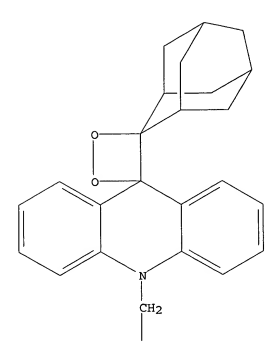
PAGE 1-A

PAGE 2-A

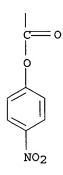
178312-97-5 HCAPLUS RN

Dispiro[acridine-9(10H),3'-[1,2]dioxetane-4',2''tricyclo[3.3.1.13,7]decane]-10-acetic acid, 4-nitrophenyl ester (9CI)
INDEX NAME) CN(CA

PAGE 1-A



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L11 ANSWER 12 OF 17 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1996:171871 HCAPLUS

DOCUMENT NUMBER: 124:225820

TITLE: Preparation of derivatized 10,10'-substituted-9,9'-

bisacridine luminescent molecules and signal solutions

INVENTOR(S): Katsilometes, George W.

PATENT ASSIGNEE(S): USA

SOURCE: PCT Int. Appl., 50 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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KIND
                            DATE
                                      APPLICATION NO.
    PATENT NO.
                                                          DATE
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                                      WO 1995-US7966
                            19960104
                                                          19950622
    WO 9600392
                      A1
       W: CN, JP, KR
       RW: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE
                      A1 19970409
                                    EP 1995-924671
                                                         19950622
       R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LI, LU, MC, NL, PT, SE
                                    CN 1995-194681
    CN 1155931
                      Α
                            19970730
                                                          19950622
    JP 10502346
                      T2
                            19980303
                                      JP 1995-503340
                                                           19950622
                                     US 1996-767288
                                                          19961216
    US 5866335
                      Α
                            19990202
                                      HK 1998-100291
                                                           19980114
    HK 1001416
                      A1
                            20050826
                                                       A 19940624
PRIORITY APPLN. INFO.:
                                      US 1994-265481
                                                    W 19950622
                                      WO 1995-US7966
```

The synthesis of 10,10'-substituted-9,9'-bisacridine mols. and their derivs. is disclosed. These mols. catalyze the production of light by chemiluminescence in the presence of a signal solution having at a pH from about 10.0 to about 14.0, at a concentration effective for producing a chemiluminescent signal, a chelating agent, a sulfoxide, a reducing sugar, and oxidant or combination of oxidants, an alc. and aqueous sodium tetraborate. These 10,10'-substituted-9,9'-biacridines are used alone or attached to haptens or macromols. and are utilized as labels in the preparation of chemiluminescent, homogeneous or heterogeneous assays. They are also used in conjunction with other chemiluminescent label mols. to produce multiple analyte chemiluminescent assays. An assay demonstrating the linearity of the signal with increasing dilns. of an anti-TSH-10,10'-paratoluo-9,9'-bisacridine conjugate is described.

IT 174569-85-8

RL: ARG (Analytical reagent use); ANST (Analytical study); USES (Uses) (preparation of bisacridine luminescent derivs. and signal solns.)

RN 174569-85-8 HCAPLUS

CN 9,9'-Biacridinium, 10,10-bis[2-[(2,5-dioxo-1-pyrrolidinyl)oxy]-2-oxoethyl]-, dinitrate (9CI) (CA INDEX NAME)

CM 1

CRN 174569-84-7 CMF C38 H28 N4 O8

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CM 2

CRN 14797-55-8

CMF N O3

L11 ANSWER 13 OF 17 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1995:599848 HCAPLUS

DOCUMENT NUMBER: 123:74302

TITLE: Netropsin and spermine conjugates of a water-soluble

quinocarcin analog: Analysis of sequence-specific DNA

interactions

AUTHOR(S): Flanagan, Mark E.; Rollins, Samuel B.; Williams,

Robert M.

CORPORATE SOURCE: Department Chemistry, Colorado State University, Ft.

Collins, CO, 80523, USA

SOURCE: Chemistry & Biology (1995), 2(3), 147-56

CODEN: CBOLE2; ISSN: 1074-5521

DOCUMENT TYPE: Journal LANGUAGE: English

Quinocarcin is the simplest of the bioxalmycin/naphthyridinomycin/tetrazom AB ine/saframycin class of antitumor antibiotics, which damage DNA in a process that is inhibited by superoxide dismutase (SOD). The oxazolidine moiety of this class of antitumor antibiotics undergoes a redox self-disproportionation reaction of the Cannizzaro type. The reaction is proposed to proceed via an intermediate carbon-centered radical, which then reduces mol. oxygen to give superoxide. We set out to determine whether the DNA-cleavage properties of these antitumor antibiotics could be retained in less complex analogs of quinocarcin. A totally synthetic, water-soluble analog of quinocarcin has been prepared This analog produced superoxide, but had considerably reduced ability to cleave supercoiled circular DNA compared to quinocarcin or tetrazomine. When conjugated to the DNA-binding mol. spermine, however, it cleaved DNA as effectively as quinocarcin at less than 1/10 the concentration A conjugate with netropsin displayed selective cleavage around the sequence 5'-d(ATTT)-3'. Mol. modeling of the interaction between the conjugate and DNA, together with the pattern of cleavage, indicates that a non-diffusable oxidant is involved in sequence-selective DNA cleavage. The spermine conjugate displayed weak antimicrobial activity. Knowledge of the stereoelectronic requirements for superoxide production by quinocarcin has allowed us to design a structurally less complex analog which has many of the same phys. properties, including water solubility, the ability to produce superoxide and the ability to cleave DNA. Covalently attaching known DNA-binding mols. to this analog gave a compound that produced sequence-specific DNA damage. Our results suggest that a mechanism other than superoxide production can mediate DNA damage by the netropsin conjugate.

IT 165253-50-9P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation of netropsin and spermine conjugates of a water-soluble quinocarcin analog and anal. of sequence-specific DNA damage)

RN 165253-50-9 HCAPLUS

CN 2-Oxa-4,10c-diazaaceanthrylene-4(1H)-acetic acid, 2a,3,5,5a,6,10b-hexahydro-10-methoxy-3,3-dimethyl-, 4-nitrophenyl ester, (2aR,5aS,10bR)-rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

L11 ANSWER 14 OF 17 HCAPLUS COPYRIGHT 2006 ACS on STN

1994:630757 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 121:230757

Preparation of quinolizinoxanthene derivatives and TITLE:

xanthene derivatives as fluorescence labeling agents

INVENTOR(S): Shiga, Masanobu

PATENT ASSIGNEE(S): Dojin Kagaku Kenkyusho Kk, Japan SOURCE:

Jpn. Kokai Tokkyo Koho, 15 pp.

CODEN: JKXXAF

DOCUMENT TYPE:

Patent Japanese LANGUAGE:

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|--------|------------|-----------------|----------|
| | | | | |
| JP 06157504 | A2 | 19940603 | JP 1992-54478 | 19920128 |
| PRIORITY APPLN. INFO.: | | | JP 1992-54478 | 19920128 |
| OTHER COHRECT(C). | MADDAT | 121.220757 | | |

OTHER SOURCE(S): MARPAT 121:230757 GI

The title compds. I [R1, R2, R9 = alkyl; or R1R8, R2R3, R9R6, or R9R7 = ring; R3,R4,R6 - R8 = H, alkyl; n = 1 - 10; Y = carboxy, etc.; X = halo] AΒ are prepared Quinolizinoxanthene II was prepared in a multiple step process starting with aminophenol derivative III. The anal. of amino acids was demonstrated using a fluorescence labeling agent of this invention. 158358-63-5P IT

RL: ARG (Analytical reagent use); SPN (Synthetic preparation); ANST (Analytical study); PREP (Preparation); USES (Uses) (preparation of quinolizinoxanthene derivs. and xanthene derivs. as fluorescence labeling agents)

RN 158358-63-5 HCAPLUS

CN 1H, 9H, 13H-Pyrido[3',2':5,6] xantheno[2,3,4-ij] quinolizinium,

7-cyano-4-[2-[(2,5-dioxo-1-pyrrolidinyl)oxy]-2-oxoethyl]-2,3,10,11,14,15-

hexahydro-5-methyl-, chloride (9CI) (CA INDEX NAME)

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L11 ANSWER 15 OF 17 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1994:435864 HCAPLUS

DOCUMENT NUMBER: 121:35864

TITLE: Fluorescent chloramphenicol derivatives for

determination of chloramphenicol acetyltransferase

activity

INVENTOR(S): Haughland, Richard P.; Kang, Hee C.; Young, Steven L.;

Melner, Michael H.

PATENT ASSIGNEE(S): Molecular Probes, Inc., USA

SOURCE: U.S., 13 pp. Cont. of U.S. Ser. No. 321,494,

abandoned.
CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

GI

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|------------------------|--------|-----------|-------------------|----------|
| | | | | |
| US 5262545 | Α | 19931116 | US 1991-722352 | 19910618 |
| US 5364764 | Α | 19941115 | US 1992-994992 | 19921221 |
| PRIORITY APPLN. INFO.: | | | US 1989-321494 B1 | 19890309 |
| | | | US 1991-722352 A3 | 19910618 |
| OTHER SOURCE(S): | MARPAT | 121:35864 | | |

OH NHCO (CH₂)_n

AB Fluorescent compds. useful in the determination of chloramphenicol

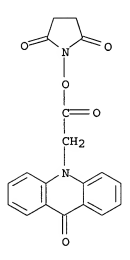
acetyltransferase (CAT) enzyme activity are described. The compds. BASE-Ns-*X are fluorescent derivs. related in structure to chloramphenicol comprising a base (I), substituted at one to five aromatic ring positions by substituents, which may be the same or different, that are alkyl, hydroxy, alkoxy, aryl, halo, nitro, amino, alkylamido, or arylamido, and 0 < n < 6; and a fluorescent moiety *X (nonreduced tricyclic difluoroboradiazaindacene fluorophore) linked to the terminal CH2 of BASE through a linker Ns (e.g., NH*X, NHCOCH2*X). The substrate compds. are acylated in the presence of CAT to produce fluorescent mono- and diacylated products, which are then phys. separated from the reaction mixture and quantitated by means of their fluorescence and/or absorbance. Fluorescent mols. conjugated to chloramphenicol include derivs. of fluorescein, rhodamine, coumarin, dimethylaminonaphthalenesulfonic acid (dansyl), pyrene, anthracene, nitrobenzoxadiazole (NBD), acridine and dipyrrometheneboron difluoride.

IT 150321-96-3

RL: RCT (Reactant); RACT (Reactant or reagent)
(fluorescent chloramphenicol derivs. for determination of chloramphenicol acetyltransferase activity)

RN 150321-96-3 HCAPLUS

CN 2,5-Pyrrolidinedione, 1-[[(9-oxo-10(9H)-acridinyl)acetyl]oxy]- (9CI) (CA INDEX NAME)



L11 ANSWER 16 OF 17 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1994:101282 HCAPLUS

DOCUMENT NUMBER: 120:101282

TITLE: Fluorescent energy transfer immunoassay

INVENTOR(S): Lakowicz, Joseph; Maliwal, Badri; Thompson, Richard;

Ozinskas, Alvydas

PATENT ASSIGNEE(S): University of Maryland, USA SOURCE: Eur. Pat. Appl., 26 pp.

CODEN: EPXXDW

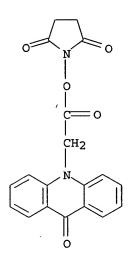
DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

| | EP 552108 | A2 | 19930721 | EP 1993-40 | 0091 | 19930115 | |
|------|----------------------|---------|-----------------|--------------|---------------|-------------|-----|
| | EP 552108 | A3 | 19930922 | | | | |
| | R: DE, FR, GB, | IT | | | | | |
| | CA 2087413 | AA | 19930718 | CA 1993-20 | 87413 | 19930115 | |
| | JP 06066802 | A2 | 19940311 | JP 1993-60 | 57 | 19930118 | |
| | JP 3325939 | B2 | 20020917 | | | | |
| | US 5631169 | Α | 19970520 | US 1994-18 | 3238 | 19940119 | |
| PRIO | RITY APPLN. INFO.: | | | US 1992-82 | 2233 A | 19920117 | |
| AB | A photoluminometric | immunc | assay compri | ses reactin | q 2 immunore | actants, 1 | |
| | labeled with a photo | | | | | | |
| | photoluminescence a | | | | | | |
| | transfer acceptor c | | | | | | |
| | radiation; and calc | | | | | | the |
| | presence of a react | | | | | | |
| | IgG labeled with the | | | | | | |
| | labeled with the ac | | | | | | |
| IT | 150321-96-3D, conju | | | | ocniocyanacc | • | |
| TI | RL: ANST (Analytica | _ | | actant | | | |
| | | | | | | | |
| | (in photoluminom | | .iiiiiunoassay) | | | | |
| RN | 150321-96-3 HCAPLU | - | | | | (007) (07 | |
| CN | 2,5-Pyrrolidinedion | e, 1-[[| (9-0xo-10(9F | ı)-acrıdınyl |) acety1]oxy] | - (9CI) (CA | |
| | INDEX NAME) | | | | | | |



L11 ANSWER 17 OF 17 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1991:535937 HCAPLUS

DOCUMENT NUMBER: 115:135937

TITLE: Preparation of N-[[(alkylideneimino)oxycarbonyl]alkyl]-

1,8-naphthalenedicarboximides and analogs as herbicide

safeners

INVENTOR(S): Saupe, Thomas; Meyer, Norbert; Plath, Peter; Schirmer,

Ulrich; Wuerzer, Bruno; Westphalen, Karl Otto; Patsch,

Manfred; Pfister, Juergen

PATENT ASSIGNEE(S): BASF A.-G., Germany

SOURCE: Eur. Pat. Appl., 45 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | | DATE |
|--------------------|----------------|------------|-----------------|---|----------|
| | | | | | |
| EP 430004 | A2 | 19910605 | EP 1990-122030 | | 19901117 |
| EP 430004 | A3 | 19911218 | | | |
| R: AT, C | CH, DE, ES, FR | R, GB, IT, | LI, NL, SE | | |
| DE 3939379 | A1 | 19910606 | DE 1989-3939379 | | 19891129 |
| DE 4021654 | A1 | 19920109 | DE 1990-4021654 | | 19900707 |
| CA 2030129 | AA | 19910530 | CA 1990-2030129 | | 19901116 |
| US 5076831 | A | 19911231 | US 1990-615865 | | 19901120 |
| JP 03190861 | A2 | 19910820 | JP 1990-323392 | | 19901128 |
| PRIORITY APPLN. IN | VFO.: | | DE 1989-3939379 | Α | 19891129 |
| | | | DE 1990-4021654 | Α | 19900707 |
| OFFICE COMPARACE. | MACICIAM | 115.12503 | 7 | | |

OTHER SOURCE(S): MARPAT 115:135937

$$R^1$$
 $Q =$
 $XCOR$
 I
 $Q =$

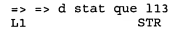
The title compds. [I; R = ON:CR5R6; R1 = 1-4 substituents which may be the same or different selected from H, halo, cyano, (halo)alkyl, etc.; R5 = H, cyano, alkyl, alkenyl, etc.; R6 = H, cyano, (halo)alkyl, alkoxy, etc.; X = (un)substituted alkylene; Y, Z = O, S] were prepared as safeners for 2-[(hetero)aryloxyphenoxy]acetate and -propionate or alkoximinomethylenecycylohexenone herbicides. Thus, I (R1 = H, X = CH2, Y = Z = O) (II); R = Cl) (preparation given) was condensed with Me2C:NOH to give II (R = ON:CMe2). II [R = ON:CR5R6; R5R6 = (CH2)3CH:C(OEt)] reduced damage to wheat of 0.03 kg/ha of the herbicide EtsCHMEH2Z1C(:NOEt)Pr (Z1 = hydroxycyclohexenonylene group Q) from 70 to 10% (with 95% control of annual ryegrass) at 0.125 kg/ha.

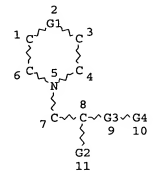
IT 135980-49-3P

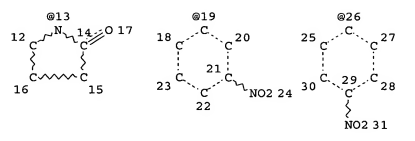
RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of, as herbicide safener)

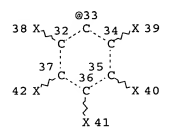
RN 135980-49-3 HCAPLUS

CN 1H-Benz[de]isoquinoline-1,3(2H)-dione, 2-[2-[(2,5-dioxo-1-pyrrolidinyl)oxy]-2-oxoethyl]- (9CI) (CA INDEX NAME)









VAR G1=C/N/O
VAR G2=O/S/N
VAR G3=O/S
VAR G4=13/19/26/33
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RING(S) ARE ISOLATED OR EMBEDDED NUMBER OF NODES IS 42

STEREO ATTRIBUTES: NONE

L5 82 SEA FILE=REGISTRY SSS FUL L1

L6 STR

VAR G1=C/N/O VAR G2=O/S/N VAR G3=O/S VAR G4=13/19/26/33 NODE ATTRIBUTES: DEFAULT MLEVEL IS ATOM DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

RSPEC 1

NUMBER OF NODES IS 42

STEREO ATTRIBUTES: NONE

46 SEA FILE=REGISTRY SUB=L5 SSS FUL L6 L7 29 SEA FILE=HCAPLUS ABB=ON PLU=ON L7 L8

36 SEA FILE=REGISTRY ABB=ON PLU=ON L5 NOT L7 L9

18 SEA FILE=HCAPLUS ABB=ON PLU=ON L9 L10

17 SEA FILE=HCAPLUS ABB=ON PLU=ON L10 NOT L8 L11

9 SEA FILE=HCAPLUS ABB=ON PLU=ON ("BARTLET JONES M"/AU OR L13 "BARTLET JONES MICHAEL"/AU) NOT (L8 OR L11)

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L13 ANSWER 1 OF 9 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2005:19284 HCAPLUS

DOCUMENT NUMBER: 142:257250

TITLE: Multiplexed protein quantitation in Saccharomyces cerevisiae using amine-reactive isobaric tagging

reagents

Ross, Philip L.; Huang, Yulin N.; Marchese, Jason N.; AUTHOR (S):

Williamson, Brian; Parker, Kenneth; Hattan, Stephen; Khainovski, Nikita; Pillai, Sasi; Dey, Subhakar; Daniels, Scott; Purkayastha, Subhasish; Juhasz, Peter;

Martin, Stephen; Bartlet-Jones, Michael; He,

Feng; Jacobson, Allan; Pappin, Darryl J.

CORPORATE SOURCE: Applied Biosystems, Framingham, MA, 01701, USA SOURCE:

Molecular and Cellular Proteomics (2004), 3(12),

1154-1169

CODEN: MCPOBS; ISSN: 1535-9476

PUBLISHER: American Society for Biochemistry and Molecular

Biology

DOCUMENT TYPE: Journal LANGUAGE: English

We describe here a multiplexed protein quantitation strategy that provides relative and absolute measurements of proteins in complex mixts. At the core of this methodol. is a multiplexed set of isobaric reagents that yield amine-derivatized peptides. The derivatized peptides are indistinguishable in MS, but exhibit intense low-mass MS/MS signature ions that support quantitation. In this study, we have examined the global protein expression of a wild-type yeast strain and the isogenic upfl Δ and xrnl Δ mutant strains that are defective in the nonsense-mediated mRNA decay and the general 5' to 3' decay pathways, resp. We also demonstrate the use of 4-fold multiplexing to enable

relative protein measurements simultaneously with determination of absolute levels of

a target protein using synthetic isobaric peptide stds. We find that inactivation of Upflp and Xrnlp causes common as well as unique effects on protein expression.

REFERENCE COUNT:

36 THERE ARE 36 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 2 OF 9 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2004:442251 HCAPLUS

DOCUMENT NUMBER:

141:238084

TITLE:

SOURCE:

Urinary N2-(2'-deoxyquanosin-8-yl)PhIP as a biomarker

for PhIP exposure

AUTHOR (S):

Fang, Min; Edwards, Robert J.; Bartlet-Jones, Michael; Taylor, Graham W.; Murray, Stephen;

Boobis, Alan R.

CORPORATE SOURCE:

Section of Experimental Medicine and Toxicology,

Imperial Coll. London, London, W12 ONN, UK Carcinogenesis (2004), 25(6), 1053-1062

CODEN: CRNGDP; ISSN: 0143-3334

PUBLISHER: Oxford University Press

DOCUMENT TYPE: Journal LANGUAGE: English

The food-derived, heterocyclic aromatic amine 2-amino-1-methyl-6-AB phenylimidazo[4,5-b]pyridine (PhIP) is genotoxic and is carcinogenic in exptl. animals. Studies on the role of PhIP in human diet-related cancer would be aided considerably by the availability of a readily applicable biomarker of the internal dose of the ultimate genotoxic species. PhIP has been shown to adduct primarily at C-8 of deoxyguanosine in DNA and so the DNA repair product N2-(2'-deoxyguanosin-8-yl)PhIP is a potential biomarker of DNA adduction and repair after exposure to PhIP. An assay for N2-(2'-deoxyguanosin-8-yl)PhIP in urine has been developed based on liquid chromatog. mass spectrometry, using a deuterated analog of the nucleoside as an internal standard and an antibody-mediated extraction procedure.

Polyclonal antibodies were raised against the PhIP-nucleotide, PhIP-nucleoside and PhIP-quanine base adducts conjugated to keyhole limpet hemocyanin. Following their evaluation, the immobilized PhIP nucleotide antibody was used for the extraction of N2-(2'-deoxyquanosin-8-vl)PhIP from The limit of detection of the assay was 125 pg and the limit of quantification 200 pg for a 50 mL human urine sample. Following oral

administration of PhIP (20 mg/kg body wt/day) to rats for 6 days, N2-(2'-deoxyguanosin-8-yl) PhIP was readily detected in the urine, reaching steady state over 3 days. This is the first direct demonstration of the urinary elimination of this adduct following exposure to parent amine. The half-life of the adduct with DNA was estimated to be .apprx.20 h. The total amount of PhIP recovered in the urine as adduct was <0.5 + 10-3% of the dose administered. Levels of the PhIP adduct in urine collections from human subjects ingesting the amine (4.9 µg) in cooked meat were below the limits of detection, indicating that humans are exposed to a bioactive dose of <3 + 10-4 of that associated with a non-carcinogenic level in rats.

REFERENCE COUNT: 61 THERE ARE 61 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 3 OF 9 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:868721 HCAPLUS

DOCUMENT NUMBER: 136:31639

TITLE: Method of stimulating non-homologous end-joining

(NHEJ) of DNA in the presence of inositol phosphate and drug screening systems and assays for compounds

that modulate NHEJ

INVENTOR(S): West, Steve Craig; Hanakahi, Leslyn Ann Akemi;

Bartlet-Jones, Michael

PATENT ASSIGNEE(S): Imperial Cancer Research Technology Limited, UK

SOURCE: PCT Int. Appl., 104 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

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APPLICATION NO.
                                                                                                                                  DATE
         PATENT NO.
                                                KIND
                                                              DATE
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                                                                                  WO 2001-GB2180
                                                                                                                                    20010518
         WO 2001090404
                20010518
W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
2408749

AB 20011129

CA 2001-GB2180

20010518
                                                  A1
                                                               20011129
                                                                                  CA 2001-2408749
EP 2001-929850
                                                               20011129
         CA 2408749
                                                   AΑ
                                                                                                                                     20010518
                                                               20030219
         EP 1283903
                                                   A1
                       AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
         JP 2004500849
                                                               20040115
                                                                                       JP 2001-586599
                                                                                                                                     20010518
                                                  T2
                                                                                                                                     20030612
                                                                                       US 2003-296014
         US 2004029130
                                                   A1
                                                               20040212
                                                                                                                               A 20000520
                                                                                       GB 2000-12179
PRIORITY APPLN. INFO.:
                                                                                       US 2000-221226P
                                                                                                                               P 20000725
                                                                                                                              P 20010214
                                                                                       US 2001-268367P
                                                                                                                              W 20010518
                                                                                       WO 2001-GB2180
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AB A method of stimulating non-homologous end-joining (NHEJ) of DNA the method comprising performing NHEJ of DNA in the presence of inositol hexakisphosphate (IP6) or other stimulatory inositol phosphate. An assay of a protein kinase wherein the assay comprises inositol hexakisphosphate (IP6) or other stimulatory inositol phosphate. The invention also provides screening assays for compds. which may modulate NHEJ and which may be therapeutically useful; and screening assays for compds. which may

modulate DNA-PK and related protein kinases and which may be therapeutically useful. Methods of modulating NHEJ and protein kinases are also disclosed. Compns. and kits that may be useful in performing the assays and methods of the invention are also claimed.

REFERENCE COUNT: 3 THERE ARE 3 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 4 OF 9 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2000:690240 HCAPLUS

DOCUMENT NUMBER: 133:330378

TITLE: Binding of inositol phosphate to DNA-PK and

stimulation of double-strand break repair

AUTHOR(S): Hanakahi, Les A.; Bartlet-Jones, Michael;

Chappell, Claire; Pappin, Darryl; West, Stephen C.

CORPORATE SOURCE: Imperial Cancer Research Fund Clare Hall Laboratories,

South Mimms, EN6 3LD, UK

SOURCE: Cell (Cambridge, Massachusetts) (2000), 102(6),

721-729

CODEN: CELLB5; ISSN: 0092-8674

PUBLISHER: Cell Press
DOCUMENT TYPE: Journal
LANGUAGE: English

AB In mammalian cells, double-strand breaks in DNA can be repaired by nonhomologous end-joining (NHEJ), a process dependent upon Ku70/80, DNA-PKCS, XRCC4, and DNA ligase IV. Starting with HeLa cell-free exts., which promote NHEJ in a reaction dependent upon all of these proteins, we have purified a novel factor that stimulates DNA end-joining in vitro. Using a combination of phosphorus NMR, mass spectroscopy, and strong anion exchange chromatog., we identify this factor as inositol hexakisphosphate (IP6). Purified IP6 is bound by DNA-PK and specifically stimulates DNA-PK-dependent end-joining in vitro. The involvement of inositol phosphate in DNA-PK-dependent NHEJ is of particular interest since the catalytic domain of DNA-PKCS is similar to that found in the phosphatidylinositol 3 (PI 3)-kinase family.

L13 ANSWER 5 OF 9 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1999:293327 HCAPLUS

DOCUMENT NUMBER: 131:128721

TITLE: Dual specificity antibodies using a double-stranded

oligonucleotide bridge

AUTHOR(S): Chaudri, Zahida N.; Bartlet-Jones, Michael;

Panayotou, George; Klonisch, Thomas; Roitt, Ivan M.;

Lund, Torben; Delves, Peter J.

CORPORATE SOURCE: Department of ImmunologyThe Windeyer Institute for

Medical Sciences, University College London, London,

W1P 6DB, UK

SOURCE: FEBS Letters (1999), 450(1,2), 23-26

CODEN: FEBLAL; ISSN: 0014-5793

PUBLISHER: Elsevier Science B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

AB The covalent conjugation of oligonucleotides to antibody Fab' fragments was optimized by using oligonucleotides modified with a hexaethylene linker arm bearing three amino groups. One oligonucleotide was coupled to antibody of one specificity and a complementary oligonucleotide to antibody of a second specificity. The antibodies were then allowed to hybridize by base pairing of the complementary nucleotide sequences and the generation of bispecific antibody was analyzed on SDS-PAGE and confirmed using BIAcore anal. The strategy of complementary oligonucleotide-linked bispecific mols. is not limited to antibodies but

is applicable to linking any two mols. of different characteristics.

THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 17

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 6 OF 9 HCAPLUS COPYRIGHT 2006 ACS on STN

1997:811809 HCAPLUS ACCESSION NUMBER:

128:154377 DOCUMENT NUMBER:

Peptide sequencing of charged derivatives by TITLE:

postsource decay MALDI mass spectrometry

Spengler, Bernhard; Luetzenkirchen, Frank; Metzger, AUTHOR (S):

Sabine; Chaurand, Pierre; Kaufmann, Raimund; Jeffery,

William; Bartlet-Jones, Michael; Pappin,

Darryl J. C.

P.O. Box 101007, Institute of Laser Medicine and CORPORATE SOURCE:

Center for Biological and Medical Research, University

of Dusseldorf, Dsseldorf, D-40001, Germany

International Journal of Mass Spectrometry and Ion SOURCE:

Processes (1997), 169/170, 127-140

CODEN: IJMPDN; ISSN: 0168-1176

Elsevier Science B.V. PUBLISHER:

DOCUMENT TYPE: Journal English LANGUAGE:

Derivatization procedures for peptides are described that can be performed with sub-picomolar amts. of sample and that are able to direct the

formation of fragment ions in Postsource Decay (PSD) MALDI mass

spectrometry. Location of a fixed charge (a quaternary ammonium ion) at the N-terminus of a peptide and modification of internal arginine residues (deletion of strong basicity) leads to a full controllability of fragment ion formation resulting in mostly complete series of N-terminal fragment The method appears to be favorably applicable to sequence anal. of unknown peptides, since in most cases the amino acid sequence can directly be read from the spectrum.

THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 26 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L13 ANSWER 7 OF 9 HCAPLUS COPYRIGHT 2006 ACS on STN

1996:236716 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 124:311580

Chemistry, mass spectrometry and peptide-mass TITLE:

databases: Evolution of methods for the rapid identification and mapping of cellular proteins

Pappin, D. J. C.; Rahman, D.; Hansen, H. F.; AUTHOR (S):

Bartlet-Jones, M.; Jeffery, W.; Bleasby, A. J.

Imperial Cancer Research Fund, London, WC2A 3PX, UK CORPORATE SOURCE: Mass Spectrometry in the Biological Sciences (1996), SOURCE:

135-50. Editor(s): Burlingame, A. L.; Carr, Steven A. Humana: Totowa, N. J.

CODEN: 62PNAY

DOCUMENT TYPE: Conference

English LANGUAGE:

Chemical, mass spectrometry, and peptide-mass databases, and methods for the identification of proteins are described,.

L13 ANSWER 8 OF 9 HCAPLUS COPYRIGHT 2006 ACS on STN

1996:159505 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 124:255185

Synthesis, evaluation and application of a panel of TITLE:

novel reagents for stepwise degradation of

polypeptides

Bures, Edward J.; Nika, Heinz; Chow, David T.; Hess, AUTHOR(S):

Daniel; Morrison, Hamish D.; Bartlet-Jones,
Michael; Pappin, Darryl J. C.; Aebersold, Ruedi
Riomedical Research Centre, University British

CORPORATE SOURCE: Biomedical Research Centre, University British

Columbia, Vancouver, Can.

SOURCE: Methods in Protein Structure Analysis, [Proceedings of

the International Conference on Methods in Protein Structure Analysis], 10th, Snowbird, Utah, Sept. 8-13, 1994 (1995), Meeting Date 1994, 57-68. Editor(s): Atassi, M. Zouhair; Appella, Ettore. Plenum: New

York, N. Y. CODEN: 62LPAK Conference

DOCUMENT TYPE: Conferent LANGUAGE: English

The authors synthesized and evaluated a panel of novel protein sequencing reagents designed to yield amino acid derivs. detectable at the low-femtomole level by electrospray-ionization mass spectrometry (ESI-MS). Protein degradation with these reagents is based on the Ph isothiocyanate functionality introduced by Edman. The chemistries were easily adapted to automated stepwise degradation Through a systematic process, the authors arrived at a new reagent, 4-(3-pyridinylmethylaminocarboxypropyl)phenyl isothiocyanate (PITC 311), that permits a sequencing approach that incorporates ESI-MS detection. By using this approach, they showed that PITC 311 is compatible with femtomole level peptide sequencing. Also mass information provided by ESI-MS detection enhances the confidence level in data interpretation. Mass information available by ESI-MS anal. of 311 PTHs assists in characterization of modified and unnatural amino acid residues. The authors aim to optimize automated sequencing cycles for high sensitivity protein sequencing, and to develop a methodol. to apply PITC 311 for high-sensitivity absorptive sequencing, and to create rapid sequencing protocols.

L13 ANSWER 9 OF 9 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1995:604181 HCAPLUS

DOCUMENT NUMBER: 123:78926

AUTHOR (S):

TITLE: The use of a volatile N-terminal degradation reagent

for rapid, high-sensitivity sequence analysis of

peptides by generation of sequence ladders

Bartlet-Jones, M.; Jeffery, W. A.; Hansen,

H. F.; Pappin, D. J. C.

CORPORATE SOURCE: Protein Sequencing Lab., Imperial Cancer Res. Fund,

London, WC2A 3PX, UK

SOURCE: Tech. Protein Chem. VI, [Pap. Symp. Protein Soc.], 8th

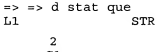
(1995), Meeting Date 1994, 3-11. Editor(s): Crabb,

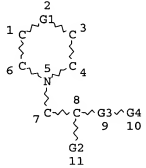
John W. Academic: San Diego, Calif.

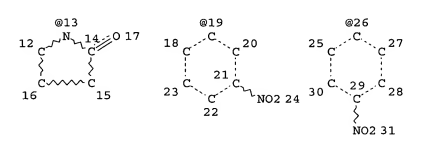
CODEN: 61MDAG

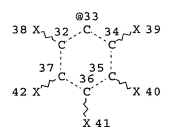
DOCUMENT TYPE: Conference LANGUAGE: English

AB The aim of this work was to explore sequencing strategies capable of rapid anal. of proteins, possibly recovered from 2-D electrophoresis gels. For this purpose, the chemical needed to be adaptable to multiple samples and sensitive enough to work in the femtomole range. The described trifluoroethyl isothiocyanate chemical is showing early signs of meeting these criteria. The demonstration, on a low-picomolar scale, that a phosphorylated tyrosine residue could be directly identified make this a potentially powerful tool for the identification of this and other sites of posttranslational modification. The inherent simplicity of the process should also allow for easy automation to permit rapid processing of samples in parallel.









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VAR G3=O/S
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DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

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STEREO ATTRIBUTES: NONE

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L6 STR

NO2 31

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GRAPH ATTRIBUTES:

RSPEC 1

NUMBER OF NODES IS 42

STEREO ATTRIBUTES: NONE

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| L12 | 103 SEA FILE=HCAPLUS ABB=ON PLU=ON ("PAPPIN D"/AU OR "PAPPIN D |
| | J"/AU OR "PAPPIN D J C"/AU OR "PAPPIN DARRYL"/AU OR "PAPPIN |
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| | CECIL"/AU OR "PAPPIN DARYL"/AU) NOT (L8 OR L11) |
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| | "BARTLET JONES MICHAEL"/AU) NOT (L8 OR L11) |
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| L15 | 92 SEA FILE=HCAPLUS ABB=ON PLU=ON L14 AND PD= <january 2004<="" 28,="" td=""></january> |
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L16 ANSWER 1 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

2004:296950 HCAPLUS

DOCUMENT NUMBER:

140:402642

TITLE:

Suppression of α -Cyano-4-hydroxycinnamic Acid

Matrix Clusters and Reduction of Chemical Noise in

MALDI-TOF Mass Spectrometry

AUTHOR(S): Smirnov, I. P.; Zhu, X.; Taylor, T.; Huang, Y.; Ross,

P.; Papayanopoulos, I. A.; Martin, S. A.; Pappin,

D. J.

CORPORATE SOURCE: Applied Biosystems, Framingham, MA, 01701, USA

SOURCE: Analytical Chemistry (2004), 76(10),

2958-2965

CODEN: ANCHAM; ISSN: 0003-2700

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal LANGUAGE: English

Progress in high-throughput MALDI-TOFMS anal., especially in proteome AB applications, requires development of practical and efficient procedures for the preparation of proteins and peptides in a form suitable for high acquisition rates. These methods should improve successful identification of peptides, which depends on the signal intensity and the absence of interfering signals. Contamination of MALDI samples with alkali salts results in reduced MALDI peptide sensitivity and causes matrix cluster formation (widely reported for CHCA matrix) observed as signals dominating in the range below m/z 1200 in MALDI spectra. One way to remove these background signals, especially for concns. of peptides lower than 10 fmol/ μ L, is to wash matrix/sample spots after peptide cocrystn. on the MALDI plate with deionized water prior to anal. This method takes advantage of the low water solubility of the CHCA compared to its alkali salts. report here that the application of some ammonium salt solns., such as citrates and phosphates, instead of deionized water greatly improves the efficiency of this washing approach. Another way to reduce matrix cluster formation is to add ammonium salts as a part of the MALDI matrix. The best results were obtained with monoammonium phosphate, which successfully suppressed matrix clusters and improved sensitivity. Combining both of these approaches-the addition of ammonium salts in the CHCA matrix followed by one postcrystn. washing step with ammonium buffer-provided a substantial (.apprx.3-5-fold) improvement in the sensitivity of MALDI-MS detection compared to unwashed sample spots. This sample preparation method resulted in improved spectral quality and was essential for successful database searching for subnanomolar concns. of protein digests.

REFERENCE COUNT: 19 THERE ARE 19 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L16 ANSWER 2 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2003:99057 HCAPLUS

DOCUMENT NUMBER: 138:151879

TITLE: Serological and proteomic evaluation of antibody

responses in the identification of tumor antigens in

renal cell carcinoma

AUTHOR(S): Unwin, Richard D.; Harnden, Patricia; Pappin,

Darryl; Rahman, Dinah; Whelan, Peter; Craven, Rachel A.; Selby, Peter J.; Banks, Rosamonde E.

CORPORATE SOURCE: Cancer Research UK Clinical Cancer Centre, St. James's

University Hospital, Leeds, LS9 7TF, UK

SOURCE: Proteomics (2003), 3(1), 45-55

CODEN: PROTC7; ISSN: 1615-9853 Wiley-VCH Verlag GmbH & Co. KGaA

DOCUMENT TYPE: Journal LANGUAGE: English

PUBLISHER:

AB Renal cell carcinoma (RCC) is relatively resistant to conventional chemotherapy and radiotherapy. However, reports of spontaneous regression along with promising results in clin. trials suggest that immunotherapuetic strategies may be of clin. benefit. Few RCC related

antiqens have been identified to date, and the tech. difficulty and time constraints of current antigen identification techniques preclude the screening of large nos. of patients. A comparatively rapid strategy has been used to identify components of tumors that elicit an antibody response in the patient - the serol. and proteomic evaluation of antibody responses (SPEAR) approach. This combines two-dimensional polyarylamide gel electrophoresis of tumor and normal kidney samples with immunoblotting using autologous patient sera and protein identification by mass spectrometry. Using the SPEAR approach to screen RCC patients for naturally occurring antitumor antibody responses, a number of candidate immunogens have been identified in patients with high-grade disease and their relative expression levels in tumor tissue compared to normal tissue have been studied. These proteins include annexins I and IV, thymidine phosphorylase (TP), carbonic anhydrase I, Mn-superoxide dismutase and major vault protein (MVP). Downstream anal. of the tissue expression of some of these proteins shows that MVP is up-regulated in 2/4 of RCC tumors but is also expressed in normal kidney whereas TP is up-regulated in 100% (11/11) of RCC cases examined with no or minimal expression in normal kidney, indicating a potential use as a therapeutic target.

REFERENCE COUNT: 43 THERE ARE 43 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L16 ANSWER 3 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2002:933437 HCAPLUS

DOCUMENT NUMBER: 139:176070

TITLE: Peptide mass fingerprinting using MALDI-TOF mass

spectrometry

AUTHOR(S): Pappin, Darryl J. C.

CORPORATE SOURCE: Imperial College, University of London, London, UK SOURCE: Methods in Molecular Biology (Totowa, NJ, United

States) (2003), 211(Protein Sequencing Protocols (2nd Edition)), 211-219 CODEN: MMBIED; ISSN: 1064-3745

PUBLISHER: Humana Press Inc.

DOCUMENT TYPE: Journal LANGUAGE: English

AB The protocols that permit reliable peptide maps to be obtained from subpicomole quantities of material using matrix-assisted laser-desorption

ionization time-of-flight (MALDI-TOF) mass spectrometry (MS) are

described. The procedures consist of staining of electroblotted proteins; enzymic digestion and elution of peptides; esterification of peptide

mixts.; MS anal. using MALDI; and database searches.

REFERENCE COUNT: 18 THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L16 ANSWER 4 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2001:724627 HCAPLUS

DOCUMENT NUMBER: 136:36207

TITLE: Changes in gene expression in macrophages infected

with Mycobacterium tuberculosis: a combined

transcriptomic and proteomic approach

AUTHOR(S): Ragno, Silvia; Romano, Maria; Howell, Steven;

Pappin, Darryl J. C.; Jenner, Peter J.;

Colston, Michael J.

CORPORATE SOURCE: Division of Mycobacterial Research, The National

Institute for Medical Research, London, NW7 1AA, UK

SOURCE: Immunology (2001), 104(1), 99-108

CODEN: IMMUAM; ISSN: 0019-2805

PUBLISHER: Blackwell Science Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

We investigated the changes which occur in gene expression in the human AB macrophage cell line, THP1, at 1, 6 and 12 h following infection with Mycobacterium tuberculosis. The anal. was carried out at the transcriptome level, using microarrays consisting of 375 human genes generally thought to be involved in immunoregulation, and at the proteomic level, using two-dimensional gel electrophoresis and mass spectrometry. The anal. of the transcriptome using microarrays revealed that many genes were up-regulated at 6 and 12 h. Most of these genes encoded proteins involved in cell migration and homing, including the chemokines interleukin (IL)-8, osteopontin, monocyte chemotactic protein-1 (MCP-1), macrophage inflammatory protein- 1α (MIP- 1α), regulated on activation, normal, T-cell expressed and secreted (RANTES), MIP-1β, MIP-3α, myeloid progenitor inhibitory factor-1 (MPIF-1), pulmonary and activation regulated chemokine (PARC), growth regulated gene- β (GRO- β), GRO- γ , MCP-2, I-309, and the T helper 2 (Th2) and eosinophil-attracting chemokine, eotaxin. Other genes involved in cell migration which were up-regulated included the matrix metalloproteinase MMP-9, vascular endothelial growth factor (VEGF) and its receptor Flk-1, the chemokine receptor CCR3, and the cell adhesion mols. vesicular cell adhesion mol.-1 (\overline{VCAM} -1) and integrin α 3. In addition to the chemokine response, genes encoding the proinflammatory cytokines IL-1β (showing a 433-fold induction), IL-2 and tumor necrosis factor- α $(TNF-\alpha)$, were also found to be induced at 6 and/or 12 h. more difficult to detect changes using the proteomic approach. Nevertheless, IL-1 β was again shown to be strongly up-regulated. enzyme manganese superoxide dismutase was also found to be strongly up-regulated; this enzyme was found to be macrophage-, rather than M. tuberculosis, derived. The heat-shock protein hsp27 was found to be down-regulated following infection. We also identified a mycobacterial protein, the product of the atpD gene (thought to be involved in the regulation of cytoplasmic pH) in the infected macrophage exts.

REFERENCE COUNT: 56 THERE ARE 56 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L16 ANSWER 5 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 2000:269897 HCAPLUS

DOCUMENT NUMBER: 133:70658

TITLE: Hydrophobic protein that copurifies with human brain

acetylcholinesterase: amino acid sequence, genomic

organization, and chromosomal localization

AUTHOR(S): Navaratnam, Dhasakumar S.; Fernando, F. Shama;

Priddle, John D.; Giles, Kurt; Clegg, Sheila M.; Pappin, Darryl J.; Craig, Ian; Smith, A. David

CORPORATE SOURCE: Department of Pharmacology, University of Oxford,

Oxford, UK

SOURCE: Journal of Neurochemistry (2000), 74(5),

2146-2153

CODEN: JONRA9; ISSN: 0022-3042
PUBLISHER: Lippincott Williams & Wilkins

DOCUMENT TYPE: Journal LANGUAGE: English

AB The mechanism of attachment of acetylcholinesterase (AChE) to neuronal membranes in interneuronal synapses is poorly understood. We have isolated, sequenced, and cloned a hydrophobic protein that co-purifies with AChE from human caudate nucleus and that we propose forms a part of a complex of membrane proteins attached to this enzyme. It is a short protein of 136 amino acids and has a mol. mass of 18 kDa. The sequence contains stretches of both hydrophobic and hydrophilic amino acids and two

cysteine residues. Anal. of the genomic sequence reveals that the coding region is divided among five short exons. Fluorescence in situ hybridization localizes the gene to chromosome 6p21.32-p21.2. Northern blot anal. shows that this gene is widely expressed in the brain

with an expression pattern that parallels that of AChE.

THERE ARE 29 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 29 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L16 ANSWER 6 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

2000:11945 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 132:148648

TITLE: Probability-based protein identification by searching

sequence databases using mass spectrometry data

Perkins, David N.; Pappin, Darryl J. C.; AUTHOR (S):

Creasy, David M.; Cottrell, John S.

CORPORATE SOURCE: Imperial Cancer Research Fund, London, WC2A 3PX, UK

SOURCE: Electrophoresis (1999), 20(18), 3551-3567

CODEN: ELCTDN; ISSN: 0173-0835

PUBLISHER: Wiley-VCH Verlag GmbH

DOCUMENT TYPE: Journal LANGUAGE: English

Several algorithms have been described in the literature for protein identification by searching a sequence database using mass spectrometry In some approaches, the exptl. data are peptide mol. wts. from the digestion of a protein by an enzyme. Other approaches use tandem mass spectrometry (MS/MS) data from one or more peptides. Still others combine mass data with amino acid sequence data. We present results from a new computer program, Mascot, which integrates all three types of search. The scoring algorithm is probability based, which has a number of advantages: (i) A simple rule can be used to judge whether a result is significant or not. This is particularly useful in guarding against false positives. Scores can be compared with those from other types of search, such as sequence homol. (iii) Search parameters can be readily optimized by iteration. The strengths and limitations of probability-based scoring are discussed, particularly in the context of high throughput, fully automated protein identification.

REFERENCE COUNT: THERE ARE 33 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L16 ANSWER 7 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1999:342725 HCAPLUS

DOCUMENT NUMBER: 131:29562

The potential use of laser capture microdissection to TITLE:

selectively obtain distinct populations of cells for

proteomic analysis. Preliminary findings

AUTHOR (S): Banks, Rosamonde E.; Dunn, Michael J.; Forbes, Mary

A.; Stanley, Anthea; Pappin, Darryl; Naven, Tom; Gough, Michael; Harnden, Patricia; Selby, Peter

ICRF Cancer Medicine Research Unit, St. James's CORPORATE SOURCE:

Hospital, Leeds, LS9 7TF, UK

Electrophoresis (1999), 20(4-5), 689-700 SOURCE:

CODEN: ELCTDN; ISSN: 0173-0835

PUBLISHER: Wiley-VCH Verlag GmbH

DOCUMENT TYPE: Journal English LANGUAGE:

Proteomics-based studies offer a powerful complementary approach to DNA/RNA-based investigations and are now being applied to investigate aspects of many diseases including cancer. The heterogeneous nature of tissue samples often makes interpretation difficult. The authors studied

the potential use of a novel laser capture microdissection (LCM) system to isolate cells of interest for subsequent proteomic anal. Retrieval of selected cells is achieved by activation of a transfer film placed in contact with a tissue section, by a laser beam (30 or 60 μm diameter) which is focused on a selected area of tissue using an inverted microscope. The precise area of film targeted by the laser bonds to the tissue beneath it and these cells are then lifted free of surrounding tissue. Although the technique was shown to be readily compatible with subsequent anal. of nucleic acids, little information is yet available regarding the application of protein-based analyses to the captured tissue. We report preliminary data regarding the potential use of the LCM system in combination with 2-D electrophoresis to examine protein profiles of selected tissue areas. Electrophoretic profiles of proteins from normal and malignant renal tissue samples showed little change following LCM, 9 selected proteins showed identical mass spectrometric sequencing profiles, and 2 selected proteins retained antigenicity. Dissection of epithelial tissue from a sample of normal human cervix resulted in enrichment of some proteins compared with anal. of the whole tissue. LCM will be a valuable adjunct to proteomic studies although further detailed validation is necessary. THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 26 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT L16 ANSWER 8 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN 1999:184612 HCAPLUS ACCESSION NUMBER: DOCUMENT NUMBER: 131:41104 Re-evaluation of the primary structure of Ralstonia TITLE: eutropha phasin and implications for polyhydroxyalkanoic acid granule binding Hanley, Steven Zachary; Pappin, Darryl J. C. AUTHOR (S): ; Rahman, Dinah; White, Andrew J.; Elborough, Kieran M.; Slabas, Antoni R. Department of Biological Sciences, University of CORPORATE SOURCE: Durham, Durham, DH13LE, UK FEBS Letters (1999), 447(1), 99-105 SOURCE: CODEN: FEBLAL; ISSN: 0014-5793 Elsevier Science B.V. PUBLISHER: Journal DOCUMENT TYPE: English LANGUAGE: Sequence anal. of several cDNAs encoding the phasin protein of Ralstonia eutropha indicated that the carboxyl terminus of the resulting derived protein sequence is different from that reported previously. This was confirmed by: (1) sequencing of the genomic DNA; (2) SDS-PAGE and peptide anal. of wild-type and recombinant phasin; and (3) mass spectrometry of wild-type phasin protein. The results have implications for the model proposed for the binding of this protein to polyhydroxyalkanoic acid granules in the bacterium. THERE ARE 27 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: 27 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT L16 ANSWER 9 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN 1997:810503 HCAPLUS

ACCESSION NUMBER:

128:100876 DOCUMENT NUMBER:

HLA-DO is a negative modulator of HLA-DM-mediated MHC TITLE:

class II peptide loading

van Ham, S. M.; Tjin, E. P. M.; Lillemeier, B. F.; AUTHOR (S):

Gruneberg, U.; van Meijgaarden, K. E.; Pastoors, L.;

Verwoerd, D.; Tulp, A.; Canas, B.; Rahman, D.;

Ottenhoff, T. H. M.; Pappin, D. J. C.;

Trowsdale, J.; Neefjes, J.

CORPORATE SOURCE:

Dep. Cellular Biochemistry, Netherlands Cancer Inst.,

Amsterdam, 1066 CX, Neth.

SOURCE:

Current Biology (1997), 7(12), 950-957 CODEN: CUBLE2; ISSN: 0960-9822

PUBLISHER:

Current Biology Ltd.

DOCUMENT TYPE:

Journal

LANGUAGE:

English

Class II mols. of the major histocompatibility complex become loaded with antigenic peptides after dissociation of invariant chain-derived peptides (CLIP) from the peptide-binding groove. The human leukocyte antigen (HLA) -DM is a prerequisite for this process, which takes place in specialized intracellular compartments. HLA-DM catalyzes the peptide-exchange process, simultaneously functioning as a peptide 'editor', favoring the presentation of stable binding peptides. Recently, HLA-DO, an unconventional class II mol., has been found associated with HLA-DM in B cells, yet its function has remained elusive. The function of the HLA-DO complex was investigated by expression of both chains of the HLA-DO heterodimer (either alone or fused to green fluorescent protein) in human Mel JuSo cells. Expression of HLA-DO resulted in greatly enhanced surface expression of CLIP via HLA-DR3, the conversion of class II complexes to the SDS-unstable phenotype and reduced antigen presentation to T-cell clones. Anal. of peptides eluted from HLA-DR3 demonstrated that CLIP was the major peptide bound to class II in the HLA-DO transfectants. Peptide exchange assays in vitro revealed that HLA-DO functions directly at the level of class II peptide loading by inhibiting the catalytic action of HLA-DM. Thus, HLA-DO is a neg. modulator of HLA-DM. By stably associating with HLA-DM, the catalytic action of HLA-DM on class II peptide loading is inhibited. HLA-DO thus affects the peptide repertoire that is eventually presented to the immune system by MHC class II mols.

REFERENCE COUNT:

THERE ARE 42 CITED REFERENCES AVAILABLE FOR THIS 42 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L16 ANSWER 10 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER:

1997:623392 HCAPLUS

DOCUMENT NUMBER:

127:306390

TITLE:

Qa-1 interaction and T cell recognition of the Qa-1

determinant modifier peptide

AUTHOR (S):

Cotterill, Lisa A.; Stauss, Hans J.; Millrain, Margaret M.; Pappin, Darryl J. C.; Rahman,

Dinah; Canas, Benito; Chandler, Phillip; Stackpoole, Arthur; Simpson, Elizabeth; Robinson, Peter J.; Dyson,

Julian P.

CORPORATE SOURCE:

Royal Postgraduate Medical School, Hammersmith

Hospital, London, W12 ONN, UK

SOURCE:

European Journal of Immunology (1997),

27(9), 2123-2132

CODEN: EJIMAF; ISSN: 0014-2980

PUBLISHER:

Wiley-VCH Journal

DOCUMENT TYPE: LANGUAGE:

English

The peptide-binding properties of the nonclassical major histocompatibility complex (MHC) class 1b mol. Qa-1 were investigated using a transfected hybrid mol. composed of the $\alpha 1$ and $\alpha 2$ domains of Qa-1b and the α 3 domain of H-2Db. This allowed the use of a monoclonal antibody directed against H-2Db while retaining the peptide-binding groove of Qa-1b. By comparison with classical MHC class I mols., intracellular maturation of the chimeric mol. was inefficient with weak intracellular association with β 2-microglobulin. However, at the cell surface the hybrid mols. were stably associated with

β2-microglobulin and were recognized by cytotoxic T lymphocyte (CTL) clones specific for the Qa-1b-presented peptide Qdm (AMAPRTLLL). A whole-cell binding assay was used to determine which residues of Qdm were important for binding to Qa-1b and CTL clones served to identify residues important for T cell recognition. Substitutions at position 1 and 5 did not reduce the efficiency of binding and had little effect on CTL recognition. In contrast, substitutions at position 9 resulted in loss of MHC class I binding. Mass spectrometric anal. of peptides eluted from immunopurified Qa-1b/ Db mols. indicated that Qdm was the dominant peptide. The closely related peptide, AMVPRTLLL, which is derived from the signal sequence of H-2Dk, was also present, although it was considerably less abundant. The mass profile suggested the presence of addnl. peptides the majority of which consisted of 8-10 amino acid residues. Finally, the finding that a peptide derived from Klebsiella pneumoniae can bind raises the possibility that this non-classical MHC class I mol. may play a role in the presentation of peptides of microorganisms.

L16 ANSWER 11 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1997:440822 HCAPLUS

DOCUMENT NUMBER: 127:201501

p47 is a cofactor for p97-mediated membrane fusion TITLE: Kondo, Hisao; Rabouille, Catherine; Newman, Richard; AUTHOR (S):

Levine, Timothy Pl; Pappin, Darryl;

Freemont, Paul; Warren, Graham

Cell Biol. Lab., Imperial Cancer Res. Fund, London, CORPORATE SOURCE:

WC2A 3PX, UK

Nature (London) (1997), 388(6637), 75-78 SOURCE:

CODEN: NATUAS; ISSN: 0028-0836

PUBLISHER: Macmillan Magazines

DOCUMENT TYPE: Journal LANGUAGE: English

At least two distinct ATPases, NSF and p97, are known to be involved in the heterotypic fusion of transport vesicles with their target membranes and the homotypic fusion of membrane compartments. The NSF-mediated fusion pathway is the best characterized, many of the components having been identified and their functions analyzed. In contrast, none of the accessory proteins for the p97-mediated fusion pathway has been identified. Now the authors have identified the first such component a protein of relative mol. mass 47,000 (p47), which forms a tight stoichiometric complex with cytosolic p97 (one trimer of p47 per hexamer of p97). It is essential for the p97-mediated regrowth of Golgi cisternae from mitotic Golgi fragments, a process restricted to animal cells. As a homolog of p47 exists in budding yeast, this indicates that it might also be involved in other membrane fusion reactions catalyzed by p97, such as karyogamy.

L16 ANSWER 12 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

1997:109668 HCAPLUS ACCESSION NUMBER:

126:222454 DOCUMENT NUMBER:

Analysis of DNA by "charge tagging" and TITLE:

matrix-assisted laser desorption/ionization mass

spectrometry

Gut, Ivo G.; Jeffery, William A.; Pappin, Darryl
J. C.; Beck, Stephan AUTHOR (S):

DNA Sequencing Laboratory, Imperial Cancer Research CORPORATE SOURCE:

Fund, London, WC2A 3PX, UK

Rapid Communications in Mass Spectrometry (SOURCE:

1997), 11(1), 43-50

CODEN: RCMSEF; ISSN: 0951-4198

PUBLISHER: Wiley
DOCUMENT TYPE: Journal
LANGUAGE: English

We have developed a method to quant. attach quaternary ammonium fixed charge tags to the 5' or 3'NH2 ends of DNA using N-hydroxysuccinimidyl ester chemical The chemical conditions for tagging were chosen so that tagging takes place exclusively on aliphatic NH2 groups while base amino groups remain unmodified. The charge tagging chemical was combined with a previously developed backbone alkylation procedure for phosphorothioate The efficiency of the detection in matrix-assisted laser desorption/ionization (MALDI) mass spectrometry of unmodified and modified DNA (phosphorothioate backbone, charge tagged, backbone alkylated, and charge tagged and backbone alkylated) was investigated using a series of different matrixes. For α -cyano-4-hydroxycinnamic acid (a matrix, commonly used for the anal. of proteins, but which gives unsatisfactory results with unmodified DNA). For instance, the charge tagged and backbone alkylated DNA is detectable with a sensitivity and resolution comparable with that for peptides. The combination of charge tagging and backbone alkylation with the use of a suitable matrix improves the detectability of small oligonucleotides by MALDI by a factor greater than 100 compared to unmodified oligonucleotides.

REFERENCE COUNT: 18 THERE ARE 18 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L16 ANSWER 13 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1997:37513 HCAPLUS

DOCUMENT NUMBER: 126:56960

TITLE: Peptide mass fingerprinting using MALDI-TOF mass

spectrometry

AUTHOR(S): Pappin, Darryl J. C.

CORPORATE SOURCE: Protein Isolation and Cloning Lab, Imperial Cancer

Research Fund, London, UK

SOURCE: Methods in Molecular Biology (Totowa, New Jersey) (

1997), 64 (Protein Sequencing Protocols),

165-173

CODEN: MMBIED; ISSN: 1064-3745

PUBLISHER: Humana
DOCUMENT TYPE: Journal
LANGUAGE: English

AB The protocols of the title method which permit reliable peptide maps to be

obtained from subpicomole quantities of material are described.

L16 ANSWER 14 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1996:398529 HCAPLUS

DOCUMENT NUMBER: 125:109439

TITLE: Identification of myocardial proteins from

two-dimensional gels by database matching of

proteolytic peptide masses

AUTHOR(S): Sutton, Chris W.; Pemberton, Kay S.; Cottrell, John

S.; Corbett, Joseph M.; Wheeler, Colin H.; Dunn,

Michael J.; Pappin, Darryl J.

CORPORATE SOURCE: Finnigan MAT Ltd., HP2 4TG, UK

SOURCE: Perspectives on Protein Engineering & Complementary

Technologies, Collected Papers, International Symposium, 3rd, Oxford, Sept. 13-17, 1994 (1995), Meeting Date 1994, 82-85. Editor(s): Geisow, Michael J.; Epton, Roger. Mayflower

Worldwide: Kingswinford, UK.

CODEN: 62ZQAP

DOCUMENT TYPE: Conference

LANGUAGE: English

Two-dimensional gels offer the most powerful method for separating complex protein mixts., but subsequent methods for analyzing individual components are slow. The identification of proteins can be accelerated by using a combination of protease digest and MALDI MS. The peptide mass spectrum of a protein represents a unique fingerprint defined by the amino acid sequence and the properties of the protease. Software has been developed so that individual peptide masses can be used to search a mass-based peptide database generated from established protein sequence databases. A list of the closest matching proteins is produced to allow identification of the sample. Examples of myocardial tissue proteins separated by 2D gel electrophoresis and identified by mass peptide fingerprinting are used to illustrate this strategy.

L16 ANSWER 15 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1996:159514 HCAPLUS

DOCUMENT NUMBER: 124:225544

TITLE: Peptide-mass fingerprinting as a tool for the rapid

identification and mapping of cellular proteins

AUTHOR(S): Pappin, D. J. C.; Rahman, D.; Hansen, H. F.;

Jeffery, W.; Sutton, C. W.

CORPORATE SOURCE: Imperial Cancer Research Fund, London, WC2A 3PX, UK

SOURCE: Methods in Protein Structure Analysis, [Proceedings of

the International Conference on Methods in Protein Structure Analysis], 10th, Snowbird, Utah, Sept. 8-13,

1994 (1995), Meeting Date 1994, 161-73.

Editor(s): Atassi, M. Zouhair; Appella, Ettore.

Plenum: New York, N. Y.

CODEN: 62LPAK Conference

DOCUMENT TYPE: Conference LANGUAGE: English

AB Simplified digestion methods based on the use of octyl glucoside that allow for the rapid, single step digestion of electro-blotted proteins in

a form suitable for both anal. by MALD spectroscopy or conventional Edman micro-sequencing are described.

L16 ANSWER 16 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1996:14825 HCAPLUS

DOCUMENT NUMBER: 124:76689

TITLE: Identification of phosphorylation sites in the mouse

estrogen receptor

AUTHOR(S): Lahooti, H.; White, R.; Hoare, S. A.; Rahman, D.;

Pappin, D. J. C.; Parker, M. G.

CORPORATE SOURCE: Mol. Endocrinology and Protein Sequencing Lab.,

Imperial Cancer Research Fund, London, WC2A 3PX, UK Journal of Steroid Biochemistry and Molecular Biology

(**1995**), 55(3/4), 305-13

CODEN: JSBBEZ; ISSN: 0960-0760

PUBLISHER: Elsevier DOCUMENT TYPE: Journal LANGUAGE: English

SOURCE:

AB Phosphorylation sites in the mouse estrogen receptor, expressed in COS-1 cells in the presence of 17β-estradiol, have been mapped by solid phase microsequencing. The receptor was first radiolabeled with [32P]orthophosphate and a number of 3H- or 14C-labeled amino acids, immunopurified and then tryptic peptides were separated by thin layer chromatog. or high performance liquid chromatog. Amino acid sequence anal. indicated that Ser-122, Ser-156, Ser-158 and Ser-298 were phosphorylated. The substitution of Ser-122 and Ser-298 with alanine had a negligible effect on the transcriptional activity of the receptor in

transfected cells. However, a reduction of transcriptional activity was observed

when Ser-122 was mutated in the context of mutations in a putative amphipathic α -helix involved in AF-2 activity. Thus, a region of AF-1 that encompasses Ser-122 appears to interact with AF-2 in the full length receptor.

L16 ANSWER 17 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1995:936888 HCAPLUS

DOCUMENT NUMBER: 124:224652

TITLE: The myristoylated alanine-rich C-kinase substrate

(MARCKS) is sequentially phosphorylated by

conventional, novel and atypical isotypes of protein

kinase C

AUTHOR(S): Herget, Thomas; Oehrlein, Silke A.; Pappin,

Darryl J. C.; Rozengurt, Enrique; Parker, Peter

.T

CORPORATE SOURCE: Institute of Physiological Chemistry, University of

Mainz, Mainz, D-55099, Germany

SOURCE: European Journal of Biochemistry (1995),

233(2), 448-57

CODEN: EJBCAI; ISSN: 0014-2956

PUBLISHER: Springer
DOCUMENT TYPE: Journal
LANGUAGE: English

The myristoylated Ala-rich C-kinase substrate (MARCKS) is the major AB protein kinase C (PKC) substrate in many cell types including fibroblasts and brain cells. The phosphorylation of MARCKS and the site specificity for different PKC isotypes are described. Conventional (c) PKC β1, novel (n) PKC δ and nPKC ϵ efficiency phosphorylated the MARCKS protein in vitro. The Km values were extremely low, reflecting a high affinity between kinases and substrate. The apparent affinity of nPKC δ (Km = 0.06 μ M) was higher than that of nPKC ϵ and cPKC β 1 (Km = 0.32 μ M). The rate of substrate phosphorylation was inversely correlated with affinity and decreased in the oder nPKC ε > cPKC β 1 > nPKC δ . Atypical (a) PKC ζ did not phosphorylate the intact MARCKS protein. However, a 25-amino-acid peptide deduced from the MARCKS phosphorylation domain, was efficiently phosphorylated by aPKC ζ as well as by the other three PKC. Site anal. revealed that only Ser residues S152, S156 and S163 were phosphorylated, with S163 phosphorylated highest, followed by S156 and S152; in contrast, S160 and S167 were not phosphorylated. No further PKC phosphorylation sites could be detected in MARCKS. The phosphorylation pattern was independent of the type of PKC isotype used. Kinetic anal. showed, that MARCKS is sequentially phosphorylated in the order S156 > S163 > S152 by cPKC, nPKC and aPKC. There was no dramatic difference in the sequential phosphorylation of MARCKS detectable when comparing the 4 PKC isotypes. The results are discussed in the context of the functional significance of MARCKS phosphorylation.

L16 ANSWER 18 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1995:491256 HCAPLUS

DOCUMENT NUMBER: 122:234807

AUTHOR(S):

TITLE: Identification of myocardial proteins from

two-dimensional gels by peptide mass fingerprinting Sutton, Chris W.; Pemberton, Kay S.; Cottrell, John

S.; Corbett, Joseph M.; Wheeler, Colin H.; Dunn,

Michael J.; Pappin, Darryl J.

CORPORATE SOURCE: Finnigan MAT Ltd., Hempstead, UK SOURCE: Electrophoresis (1995), 16(3), 308-16

CODEN: ELCTDN; ISSN: 0173-0835

PUBLISHER: VCH

DOCUMENT TYPE: Journal

LANGUAGE: English

Two-dimensional gels offer a powerful method for separating complex protein mixts., but subsequent methods for analyzing individual components, such as protein sequencing and Western immunoblotting, are laborious and slow. The identification of proteins can be accelerated by using a combination of protease digestion and matrix assisted laser desorption-mass spectrometry (MALDI-MS). The peptide mass spectrum of a protein represents a unique fingerprint determined by the amino acid sequence and the cleavage properties of the protease. Software has been developed so that peptide masses can be used to search a mass-based peptide database generated from established protein sequence databases. A list of the closest matching proteins is produced to allow identification of the The strategy was applied to 52 protein spots from human myocardial tissue separated by two-dimensional electrophoresis (2-DE) gels and analyzed blind. Conditions for optimal trypsin digestion of proteins electroblotted onto polyvinylidene difluoride (PVDF) membranes are described. Mass data were generated from both Coomassie Brilliant Blue and sulforhodamine B-stained proteins, though the former required destaining prior to digestion. Alkylation of cysteine and oxidation of methionine were significant modifications that influenced the successful identification of a protein spot. Examples are presented to illustrate the advantages and disadvantages of this approach.

L16 ANSWER 19 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1994:675783 HCAPLUS

DOCUMENT NUMBER: 121:275783

TITLE: Peptide ladder sequencing by mass spectrometry using a

novel, volatile degradation reagent

AUTHOR(S): Jones, Michael Bartlet; Jeffrey, William A.; Hansen,

Hans F.; Pappin, Darryl J. C.

CORPORATE SOURCE: Protein Sequencing Lab., Imp. Cancer Res. Fund,

London, WC2A 3PX, UK

SOURCE: Rapid Communications in Mass Spectrometry (

1994), 8(9), 737-42

CODEN: RCMSEF; ISSN: 0951-4198

DOCUMENT TYPE: Journal LANGUAGE: English

A conceptually novel approach to protein sequencing involves the generation of ragged-end polypeptide chains followed by mass spectroscopic anal. of the resulting nested set of fragments. We report here on the synthesis and development of a volatile isothiocyanate (trifluoroethylisothiocyanate) that allows the identification of several consecutive residues starting with a few picomoles of peptide. The nested set of peptides is generated simply by adding equal aliquots of starting peptide each cycle and driving both the coupling and cleavage reactions to completion. No addnl. reagents are required to act as chain terminators and retention of the peptide terminal amine allows for subsequent modification with quaternary ammonium alkyl NHS esters to improve sensitivity. Complex washing procedures are not required each cycle, as reagents and byproducts are efficiently removed under vacuum, eliminating extractive loss. Multiple peptide samples can be proposed simultaneously, with each degradation cycle completed in 35-40 min. The inherent simplicity of the process should allow for easy automation and permit rapid processing of samples in parallel.

L16 ANSWER 20 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1994:647959 HCAPLUS

DOCUMENT NUMBER: 121:247959

Competition between nuclear localization and secretory TITLE:

signals determines the subcellular fate of a single

CUG-initiated form of FGF3

Kiefers, Paul; Acland, Piers; Pappin, Darryl AUTHOR (S):

; Peters, Gordon; Dickson, Clive

CORPORATE SOURCE: Imperial Cancer Research Lund, London, WC2A 3PX, UK

SOURCE: EMBO Journal (1994), 13(17), 4126-36

CODEN: EMJODG; ISSN: 0261-4189

Journal DOCUMENT TYPE: LANGUAGE: English

The presumed open reading frame for mouse FGF3, starting at the most 5' AUG codon, predicts a hydrophobic N-terminus characteristic of a signal peptide for secretion. However, in reticulocyte lysates and transfected COS-1 cells, the full-length Fgf-3 cDNA is translated almost exclusively from an upstream CUG codon. The resultant products are distributed in both the nucleus and the secretory pathway, implying that the single CUG-initiated form of FGF3 has dual fates. By analyzing a series of deletion and replacement mutants and by linking parts of FGF3 to a heterologous protein, we show that secretion is mediated by cleavage adjacent to the previously defined signal peptide, whereas nuclear localization is determined primarily by a classical but relatively weak bipartite motif. In the context of FGF3, nuclear localization also requires the N-terminal sequences which lie upstream of the signal peptide. Thus, the subcellular fate of FGF3 is determined by the competing effects of signals for secretion and nuclear localization within the same protein, rather than by alternative initiation or processing.

L16 ANSWER 21 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

1994:75401 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 120:75401

TITLE: Tyrosine phosphorylation of α tubulin in human T

lymphocytes

AUTHOR (S): Ley, Steven C.; Verbi, Winston; Pappin, Darryl J.

C.; Druker, Brian; Davies, Adelina A.; Crumpton,

Michael J.

Natl. Inst. Med. Res., London, UK CORPORATE SOURCE:

European Journal of Immunology (1994), SOURCE:

24(1), 99-106 CODEN: EJIMAF; ISSN: 0014-2980

DOCUMENT TYPE: Journal LANGUAGE: English

N-terminal sequencing of the 55- and 50-kDa polypeptides affinity purified

on a phosphotyrosine monoclonal antibody column from activated Jurkat T cells identified α and β tubulin. Two-dimensional gel anal. indicated that α tubulin was directly phosphorylated

on tyrosine. β Tubulin was not detectably tyrosine phosphorylated

but was precipitated by anti-phosphotyrosine (PTyr) antibody by virtue of its association with the α subunit as a heterodimer. Phosphotyrosyl α

tubulin was not incorporated into intact microtubules and was all in the

unpolymd. soluble fraction. Thus, tyrosine phosphorylation of α tubulin may inhibit the ability of this subunit to polymerize into

microtubules. Stimulation of Jurkat T cells via T cell receptor increased the amount of tubulin precipitated by the anti-PTyr antibody. These data

possibility that the polymerization of tubulin heterodimers may be regulated by phosphorylation on tyrosine during T cell activation.

L16 ANSWER 22 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN ACCESSION NUMBER: 1993:644768 HCAPLUS

119:244768 DOCUMENT NUMBER:

Rapid indentification of proteins by peptide-mass TITLE:

fingerprinting. [Erratum to document cited in

CA119(15):155221x]

Pappin, D. J.; Hojrup, P.; Bleasby, A. J. AUTHOR(S):

Protein Sequencing Lab., Imp. Cancer Res. Fund, CORPORATE SOURCE:

London, WC2A 3PX, UK

Current Biology (1993), 3(7), 487 SOURCE:

CODEN: CUBLE2; ISSN: 0960-9822

Journal DOCUMENT TYPE: English LANGUAGE:

The errors were not reflected in the abstract or the index entries. AB

L16 ANSWER 23 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

1993:555221 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 119:155221

Rapid indentification of proteins by peptide-mass TITLE:

fingerprinting

Pappin, D. J. C.; Hojrup, P.; Bleasby, A. J. AUTHOR(S): CORPORATE SOURCE: Protein Sequencing Lab., Imp. Cancer Res. Fund,

London, WC2A 3PX, UK

Current Biology (1993), 3(6), 327-32 CODEN: CUBLE2; ISSN: 0960-9822 SOURCE:

DOCUMENT TYPE: Journal English LANGUAGE:

The authors report the development of the mol. weight search (MOWSE) peptide-mass database at the SERC Daresbury Laboratory Practical experience showed that sample proteins can be identified uniquely from as few as 3 or 4 exptl. determined peptide masses when these are screened against a fragment database that is derived from >50,000 proteins. Peptide-mass fingerprints can prove as discriminating as linear peptide sequences but can be obtained in a fraction of the time using less protein. In many cases, this allows for a rapid identification of a sample protein before committing it to protein sequence anal. Fragment masses also provide information, at the protein level, that is complementary to the information provided by large-scale DNA sequencing or mapping projects.

L16 ANSWER 24 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

1991:581148 HCAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 115:181148

TITLE: Purification and characterization of biologically active scatter factor from ras-transformed NIH 3T3

conditioned medium

Coffer, Arnold; Fellows, Jane; Young, Susan; AUTHOR(S):

Pappin, Darryl; Rahman, Dinah

Protein Isol. Cloning Lab., Imp. Cancer Res. Fund, CORPORATE SOURCE:

London, WC2A 3PX, UK

SOURCE: Biochemical Journal (1991), 278(1), 35-41

CODEN: BIJOAK; ISSN: 0306-3275

DOCUMENT TYPE: Journal LANGUAGE: English

Scatter factor (SF), a glycoprotein produced by cultured fibroblasts, acts in vitro on epithelial cells causing separation and increased local motility. In this study, the polypeptide was purified to apparent homogeneity in high yields with conserved biol. activity from medium conditioned by ras-transformed NIH 3T3 cells, by a 3-step procedure involving ammonium sulfate fractionation, cation-exchange, and hydroxyapatite chromatog. After purification, SF specific activity increased from .apprx.0.3 units/µg in unprocessed conditioned medium to .apprx.5 units/ng, and cumulative recovery of biol. activity was .apprx.38%. Treatment of pure SF with

N-qlycanase resulted in a decreased Mr, but no concomitant effect was observed on biol. activity. Proteolytic activity was absent from samples of both partially purified and pure SF. The biochem. studies showed that SF, which is highly aggregated in low-ionic-strength media, is not aggregated in 0.4 M-salt. Under non-reducing conditions, pure SF migrated as a single strained band at Mr 67,000 on SDS/PAGE, and biol. activity was eluted from unstained gels with an identical Mr. SF was electrofocused sharply at pI 8.5 with no degradation of activity. From ultracentrifugation studies (under non-aggregating conditions), the sedimentation coefficient of active SF was 3.7S and f.p.l.c. mol. sieve chromatog. indicated a Stokes' radius of 2.95 nm. The calculated Mr from these data was 61,400. The appearance of 3 stained polypeptides of Mr 82,000, 57,000, and 32,000 derived from the Mr-67,000 constituent after reduction with mercaptoethanol suggests that SF may be a heterodimer of Mr-57,000 and -32,000 subunits. Data from protein sequence anal. of the hydroxyapatite-purified protein confirms that SF has sequence identity with both rat hepatocyte growth factor and human fibroblast tumor cytotoxic factor.

L16 ANSWER 25 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1991:467672 HCAPLUS

DOCUMENT NUMBER: 115:67672

TITLE: New approaches to covalent sequence analysis

Pappin, Darryl J. C.; Coull, James M.; AUTHOR (S):

Koester, Hubert

CORPORATE SOURCE: MilliGen/Biosearch Div., Millipore, Burlington, MA,

01803, USA

SOURCE: Curr. Res. Protein Chem.: Tech., Struct., Funct.,

[Pap. Annu. Symp. Protein Soc.], 3rd (1990),

Meeting Date 1989, 191-202. Editor(s): Villafranca,

Joseph J. Academic: San Diego, Calif.

CODEN: 56XQAW

DOCUMENT TYPE: Conference

LANGUAGE: English

A symposium report on covalent (solid-phase) sequence anal. of proteins. Thus, peptides or proteins are blotted onto an underivatized polyvinylidene membranes, stained by conventional techniques, and then efficiently covalently immobilized to the membrane surface by entrapment in a thin polymer coating.

L16 ANSWER 26 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1991:425568 HCAPLUS

DOCUMENT NUMBER: 115:25568

TITLE: Immobilization of proteins and peptides on insoluble

supports for sequencing and other applications

INVENTOR(S): Pappin, Darryl J. C.; Coull, James M.;

Koester, Hubert

PATENT ASSIGNEE(S): Millipore Corp., USA

SOURCE: Eur. Pat. Appl., 18 pp.

CODEN: EPXXDW

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

| PATENT NO. | KIND | DATE | APPLICATION NO. | DATE |
|----------------|----------|----------|-----------------|------------|
| EP 410323 | A2 | 19910130 | EP 1990-113972 | 19900720 < |
| EP 410323 | A2 A3 | 19910130 | EP 1990-1139/2 | 19900720 < |
| R: DE, FR, GB, | IT, NL | , SE | | |
| US 5071909 | Α | 19911210 | US 1989-385711 | 19890726 < |

JP 03141300 A2 19910617 JP 1990-194113 19900724 <-PRIORITY APPLN. INFO.: US 1989-385711 A 19890726

AB A peptide or protein is immobilized onto a flat, microporous membrane by (1) adsorbing the peptide or protein and a crosslinkable polymer onto the membrane surface, and (2) crosslinking the polymer to produce a polymer network entrapping the protein or peptide therein. The immobilized peptide or protein is suitable for sequence anal. or other chemical or enzymic processes. Thus, a polyvinylidene difluoride membrane disk containing electroblotted β-lactoglobulin A and stained with sulforhodamine B was treated with diisopropyl-carbodiimide and methylenedianiline (polymer crosslinking agent), dried, then treated with polyacrylic acid (5000 mol. weight). The prepared disk was subjected to 20 cycles of Edman degradation The initial sequencing yield was 35 pmol and the repetitive yield 90%.

L16 ANSWER 27 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1991:243669 HCAPLUS

DOCUMENT NUMBER: 114:243669

TITLE: Functionalized membrane supports for covalent protein

microsequence analysis

AUTHOR(S): Coull, James M.; Pappin, Darryl J. C.; Mark,

Jonathan; Aebersold, Ruedi; Koster, Hubert

CORPORATE SOURCE: MilliGen/Bios., Div. Millipore, Burlington, MA, 01803,

USA

SOURCE: Analytical Biochemistry (1991), 194(1),

110-20

CODEN: ANBCA2; ISSN: 0003-2697

DOCUMENT TYPE: Journal LANGUAGE: English

Methods were developed for high-yield covalent attachment of peptides and proteins to isothiocyanate and arylamine-derivatized poly(vinylidene difluoride) membranes for solid-phase sequence anal. Solns. of protein or peptide were dried onto 8-mm membrane disks such that the functional groups on the surface and the polypeptide were brought into close proximity. In the case of the isothiocyanate membrane, reaction between polypeptide amino groups and the surface isothiocyanate moieties was promoted by application of aqueous N-methylmorpholine. Attachment of proteins and peptides to the arylamine surface was achieved by application of water-soluble carbodiimide in a pH 5.0 buffer. Edman degradation of covalently bound polypeptides was accomplished with initial and repetitive sequence yields ranging 33-75% and 88.5-98.5%, resp. The yields were independent of the sample load (20 pmol to >1 nmol) for either surface. Significant loss of material was not observed when attachment residues were encountered during sequence runs. Application of bovine β -lactoglobulin A chain, staphylococcus protein A, or the peptide melittin to the isothiocyanate membrane allowed for extended N-terminal sequence identification (35 residues from 20 pmol of $\beta\text{--}$ lactoglobulin). Several synthetic and naturally occurring peptides were sequenced to the C-terminal residue following attachment to the arylamine In 1 example, 10 μg of bovine α -casein was digested with staphylococcal protease V8 and the peptides were separated by reversed-phase chromatog. Peptide fractions were then directly applied to arylamine membrane disks for covalent sequence anal. From as little as 2 pmol of initial signal it was possible to determine substantial sequence information (>10 residues).

L16 ANSWER 28 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1990:420480 HCAPLUS

DOCUMENT NUMBER: 113:20480

TITLE: Solid-phase sequence analysis of proteins

electroblotted or spotted onto polyvinylidene

difluoride membranes

AUTHOR(S): Pappin, Darryl J. C.; Coull, James M.;

Koster, Hubert

CORPORATE SOURCE: MilliGen/Biosearch, Burlington, MA, 01803, USA

SOURCE: Analytical Biochemistry (1990), 187(1),

10-19

CODEN: ANBCA2; ISSN: 0003-2697

DOCUMENT TYPE: Journal LANGUAGE: English

Electroblotted proteins noncovalently bound to polyvinylidene difluoride (PVDF) membranes are typically sequenced using adsorptive sequencer protocols (gas phase or pulsed-liquid) that do not require a covalent linkage between protein and surface. Simple chemical protocols were developed where proteins are first electroblotted onto unmodified PVDF membranes, visualized with common protein stains, and then immobilized for solid-phase sequence anal. Adsorbed, stained proteins are first treated with phenylisothiocyanate (PITC) to modify α and ϵ amines. The protein is then overlayed with a solution of 1,4-phenylene diisothiocyanate (DITC), followed by a few microliters of a basic solution containing a poly(alkylamine). As the polymer dries onto the surface both polymer and remaining protein amino groups are crosslinked by DITC. The protein is thus immobilized to the membrane surface by entrapment in a thin polymer coating. The coating is transparent to the degradation chemical, and extensive enough to remain immobilized even in the absence of any covalent link between polymer and surface. Partial modification with PITC allows for identification of N-terminal and internal lysine residues during sequencing. The process was tested with a variety of poly(alkylamines), linear and branched, with mol. wts. ranging from 600 to >100,000. Proteins bound in this manner were successfully sequenced using covalent (solid-phase) sequencer protocols with cyclic times as short as 26 min.

L16 ANSWER 29 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1990:113201 HCAPLUS

DOCUMENT NUMBER: 112:113201

TITLE: The extrinsic 33 kDa polypeptide of the

oxygen-evolving complex of photosystem II is a putative calcium-binding protein and is encoded by a

multi-gene family in pea

AUTHOR(S): Wales, Richard; Newman, Barbara J.; Pappin,

Darryl; Gray, John C.

CORPORATE SOURCE: Bot. Sch., Univ. Cambridge, Cambridge, CB2 3EA, UK

SOURCE: Plant Molecular Biology (1989), 12(4),

439-51

CODEN: PMBIDB; ISSN: 0167-4412

DOCUMENT TYPE: Journal LANGUAGE: English

AB The extrinsic 33 kDa polypeptide of the water-oxidizing complex has been extracted from pea photosystem II particles by washing with alkaline-Tris and purified by ion-exchange chromatog. The N-terminal amino acid sequence has been determined, and specific antisera have been raised in rabbits and used to screen a pea leaf cDNA library in λgt 11. Determination of the nucleotide sequence of pos. clones revealed an essentially full-length cDNA for the 33 kDa polypeptide, the deduced amino acid sequence showing it to code for a mature protein of 248 amino acids with an N-terminal transit peptide of 81 amino acids. The protein showed a high degree of conservation with previously reported sequences for the 33 kDa protein from other species and the sequence contained a putative Ca2+-binding site with homol. to mammalian intestinal calcium-binding proteins. Northern

anal. of total pea RNA indicated a message of approx. 1.4 kb, in good agreement with the size of the cDNA obtained at indicated a message of approx. 1.4 kb, in good agreement with the size of the cDNA obtained at 1.3 kbp. Southern blots of genomic DNA probed with the labeled cDNA give rise to several bands suggesting that the 33 kDa polypeptide is coded by a multi-gene family.

L16 ANSWER 30 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1989:611670 HCAPLUS

111:211670 DOCUMENT NUMBER:

Exported proteins of Neurospora crassa: TITLE:

1-glucoamylase

Koh-Luar, Siok Im; Parish, J. H.; Bleasby, A. J.; AUTHOR (S):

Pappin, D. J. C.; Ainley, K.; Johansen, F. E.;

McPherson, M. J.; Radford, A.

Dep. Biochem., Univ. Leeds, Leeds, LS2 9JT, UK CORPORATE SOURCE:

Enzyme and Microbial Technology (1989), SOURCE:

11(10), 692-5

CODEN: EMTED2; ISSN: 0141-0229

DOCUMENT TYPE: Journal LANGUAGE: English

Polypeptides were extracted from the culture supernatants of N. crassa growing in a variety of media. The polypeptides were analyzed by quant. SDS-PAGE and the single polypeptide produced in the largest amount under the conditions tested was of mol. weight 69,000 and was referred to as p69. synthesis of p69 was induced by starch and maltose, and studies on a partially purified preparation established that it was a glucoamylase. sequence of the N-terminal 30 amino acids of p69 showed that this was related to sequences of amylases from Aspergillus species. A synthetic oligonucleotide probe was synthesized and used to search for the corresponding gene(s) in Neurospora. A unique region was identified and mapped in a clone from a genomic library. The implications of the work for the development of Neurospora export and expression systems are discussed.

L16 ANSWER 31 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1988:625726 HCAPLUS

DOCUMENT NUMBER: 109:225726

Amino acid sequence analysis of the small TITLE:

subunit of ribulose bisphosphate carboxylase from

Fucus (Phaeophyceae)

Keen, Jeffrey N.; Pappin, Darryl J. C.; AUTHOR (S):

Evans, Leonard V.

Dep. Pure Appl. Biol., Univ. Leeds, Leeds, LS2 9JT, UK CORPORATE SOURCE:

Journal of Phycology (1988), 24(3), 324-7 CODEN: JPYLAJ; ISSN: 0022-3646 SOURCE:

DOCUMENT TYPE: Journal English LANGUAGE:

Amino acid sequence information for the small subunit of ribulose 1,5-bisphosphate carboxylase/oxygenase (Rubisco) from a nongreen alga was reported. N-terminal sequences are presented for the polypeptide from three species of the genus Fucus (Phaeophyceae). Although homologous to small subunit polypeptides from other organisms, the Fucus sequences exhibit a unique N-terminal section resembling neither cyanobacterial nor chlorophytic sequences. This difference may be a consequence of the plastid DNA coding arrangement for the small subunit in chromophytes, a situation reported for the related organism Olisthodiscus but not previously investigated at the amino acid sequence level.

L16 ANSWER 32 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1988:182407 HCAPLUS

DOCUMENT NUMBER: 108:182407

TITLE: Isolation and characterization of a minor legumin and

its constituent polypeptides from Pisum sativum (pea)

AUTHOR(S): March, John F.; Pappin, Darryl J. C.; Casey,

Rod

CORPORATE SOURCE: John Innes Inst., Norwich, NR4 7UH, UK

SOURCE: Biochemical Journal (1988), 250(3), 911-15

CODEN: BIJOAK; ISSN: 0306-3275

DOCUMENT TYPE: Journal LANGUAGE: English

AB The purification and characterization of a minor legumin species from P. sativum is described. Electrophoretic data indicate that it corresponds

to a legumin subunit pair previously designated L1. The

β-polypeptides of the minor legumin have a phenylalanine N-terminus. This is the first time that an amino acid other than glycine has been reported as the N-terminus of the basic polypeptides from legumin-like

proteins from any plant species. Sequence analyses of the isolated α - and β -polypeptides of the minor legumin show that

it does not correspond to any of the 3 legumin gene families previously

defined on the basis of DNA hybridiations and genetic analyses.

L16 ANSWER 33 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1988:147347 HCAPLUS

DOCUMENT NUMBER: 108:147347

TITLE: N-terminal amino acid sequence analysis of

the subunits of pea photosystem I

AUTHOR(S): Dunn, Paul P. J.; Packman, Leonard C.; Pappin,

Darryl; Gray, John C.

CORPORATE SOURCE: Dep. Bot., Univ. Cambridge, Cambridge, CB2 1QW, UK

SOURCE: FEBS Letters (1988), 228(1), 157-61

CODEN: FEBLAL; ISSN: 0014-5793

DOCUMENT TYPE: Journal LANGUAGE: English

AB Six core subunits of pea photosystem I were isolated and their N-terminal amino acid sequences determined by gas-phase or solid-phase sequencing. On average

>30 residues were determined from the N-terminus of each polypeptide. Sequence anal. revealed three polypeptides with charged N-terminal regions (21, 17 and 11 kDa subunits), one polypeptide with a predominantly hydrophobic N-terminal region (9 kDa subunit), one polypeptide which is cysteine-rich (8 kDa subunit) and one which is alanine-rich (13 kDa subunit).

L16 ANSWER 34 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1987:191404 HCAPLUS

DOCUMENT NUMBER: 106:191404

TITLE: Homology between the pyrazine-binding protein from

nasal mucosa and major urinary proteins

AUTHOR(S): Cavaggioni, A.; Sorbi, R. T.; Keen, J. N.;

Pappin, D. J. C.; Findlay, J. B. C.

CORPORATE SOURCE: Ist. Fisiol., Univ. Parma, Parma, 43100, India

SOURCE: FEBS Letters (1987), 212(2), 225-8

CODEN: FEBLAL; ISSN: 0014-5793

DOCUMENT TYPE: Journal LANGUAGE: English

AB Sequence anal. of the pyrazine-binding protein from bovine olfactory mucosa reveals marked homol. with a family of proteins of unknown function found in the urine of the adult male mouse and rat. In view of the dramatic biol. responses to odorants transmitted in male

rodent urines, it is proposed that these proteins play important roles in some aspects of odor transmission and reception.

L16 ANSWER 35 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1986:203984 HCAPLUS

DOCUMENT NUMBER: 104:203984

TITLE: Posttranslational processing of concanavalin A

precursors in jackbean cotyledons

AUTHOR(S): Bowles, Dianna J.; Marcus, Susan E.; Pappin,

Darryl J. C.; Findlay, John B. C.; Eliopoulos,

Elias; Maycox, Peter R.; Burgess, Jeremy

CORPORATE SOURCE: Dep. Biochem., Univ. Leeds, Leeds, LS2 9JT, UK

SOURCE: Journal of Cell Biology (1986), 102(4),

1284-97

CODEN: JCLBA3; ISSN: 0021-9525

DOCUMENT TYPE: Journal LANGUAGE: English

Metabolic labeling of immature jackbean cotyledons with 14C-amino acids was used to determine the processing steps involved in the assembly of concanavalin A. Pulse-chase expts. and analyses of immunopptd. lectin forms indicated a complex series of events involving 7 distinct species. The structural relatedness of all of the intermediate species was confirmed by 2-dimensional mapping of 125I-tryptic peptides. An initial qlycosylated precursor was deglycosylated and cleaved into smaller polypeptides, which subsequently reannealed over a period of 10-27 h. NH2-terminal sequencing of the abundant precursors confirmed that the intact subunit of concanavalin A was formed by the reannealing of 2 fragments, since the alignment of residues 1-118 and 119-237 was reversed in the final form of the lectin identified in the chase and the precursor first labeled. When the tissue was pulse-chased in the presence of monensin, processing of the glycosylated precursor was inhibited. weak bases NH4Cl and chloroquine were without effect. Immunocytochem. studies showed that monensin treatment caused the accumulation of immunoreactive material at the cell surface and indicated that the ionophore had induced the secretion of a component normally destined for deposition within the protein bodies. Consideration of the tertiary structure of the glycosylated precursor and mature lectin showed that the entire series of processing events could occur without significant refolding of the initial translational product. Proteolytic events included removal of a peptide from the surface of the precursor mol. that connected the NH2- and COOH-termini of the mature protein. This processing activated the carbohydrate-binding activity of the lectin. chase data suggested the occurrence of a simultaneous cleavage and formation of a peptide bond, raising the possibility that annealment of the fragments to give rise to the mature subunit involves a transpeptidation event rather than cleavage and subsequent religation.

L16 ANSWER 36 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1986:164913 HCAPLUS

DOCUMENT NUMBER: 104:164913

TITLE: Coronavirus IBV glycopolypeptides: locational studies using proteases and saponin, a membrane permeabilizer

AUTHOR(S): Cavanagh, David; Davis, Philip J.; Pappin, Darryl

J. C.

CORPORATE SOURCE: Houghton Poultry Res. Statn.,

Houghton/Huntingdon/Cambs., PE17 2DA, UK

SOURCE: Virus Research (1986), 4(2), 145-56

CODEN: VIREDF; ISSN: 0168-1702

DOCUMENT TYPE: Journal LANGUAGE: English

[35S] methionine-labeled avian infectious bronchitis virus (IBV) (strain AB 41) and its purified protein components and virions of IBV-Beaudette were incubated with 10 proteases. Several proteases hydrolyzed all or some of the membrane glycopolypeptide M (mol. weight 30 kilodaltons [30K]) and removed .apprx.1.3K of peptide from the N terminus plus both glycans, as determined by SDS-polyacrylamide gel electrophoresis. N-terminal anal . of [3H]isoleucine-labeled M after hydrolysis by bromelain revealed that the first 9 residues were removed. After the virions were permeabilized with saponin, a further 2.5K decrease in mol. weight was produced from the C terminus. When considered with the hydropathicity plot anal. of the amino acid sequence of M, as few as 9-20 N-terminal amino acids may protrude at the outer membrane surface, and that there msy be a highly protease sensitive sequence of .apprx.20-25 residues at the C-terminus of M exposed in the lumen of the virion. S2 but not S1 was cleaved to a major glycopolypeptide of .apprx.71K by several proteases, and to 76K by trypsin. N-terminal sequencing of the 71K glycopolypeptide revealed that it had the same N terminus as intact S2. After hydrolysis in the presence and absence of saponin, it was concluded that S2 is very sensitive to hydrolysis near its C terminus at residues close to the outer membrane surface.

L16 ANSWER 37 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1985:590791 HCAPLUS

DOCUMENT NUMBER: 103:190791

TITLE: The primary sequence of Ricinus communis agglutinin.

Comparison with ricin

AUTHOR(S): Roberts, Lynne M.; Lamb, F. Ian; Pappin, Darryl

J. C.; Lord, J. Michael

CORPORATE SOURCE: Dep. Biol. Sci., University of Warwick, Coventry, CV4

7AL, UK

SOURCE: Journal of Biological Chemistry (1985),

260(29), 15682-6

CODEN: JBCHA3; ISSN: 0021-9258

DOCUMENT TYPE: Journal LANGUAGE: English

A mixture of synthetic oligonucleotides representing all possible sequences of a peptide present in the ricin B chain was used to screen a cDNA library constructed using ripening castor bean seed poly(A+) RNA. The 8 largest recombinant plasmids selected, by hybridization, a single mRNAspecies that encoded preprolectin, as shown by immunopptn. enzyme anal. of these clones showed 2 classes of sequences complementary to 2 distinct, but closely related, preprolectin mRNA species. The nucleotide sequence of the cloned cDNA from 1 of these classes encodes preproricin and has been presented elsewhere. nucleotide sequence of the 2nd class is presented and represents the preproagglutinin of R. communis. The entire coding sequence was deduced from 2 overlapping cDNA clones with inserts of 1668 and 1151 base pairs. The coding region defines a preproprotein with a 24-amino acid N-terminal signal sequence preceding the A chain (266 amino acids) which is joined to the B chain (262 amino acids) by a 12-amino acid linking peptide. The protein was confirmed as R. communis agglutinin since the deduced B chain N-terminal sequence corresponds exactly with that determined for purified R. communis agglutinin B chain over a region where several residue differences occur in the ricin B chain. The nucleotide and deduced amino acid sequences of the R. communis agglutinin precursor are compared with those of the ricin precursor.

L16 ANSWER 38 OF 38 HCAPLUS COPYRIGHT 2006 ACS on STN

ACCESSION NUMBER: 1985:401514 HCAPLUS

DOCUMENT NUMBER: 103:1514

TITLE: Cloning and sequencing of the gene encoding the spike

protein of the coronavirus IBV

AUTHOR(S): Binns, Matthew M.; Boursnell, Michael E. G.; Cavanagh,

David; Pappin, Darryl J. C.; Brown, T. David

Κ.

CORPORATE SOURCE: Houghton Poultry Res. Stn.,

Houghton/Huntingdon/Cambs., PE17 2DA, UK
Journal of General Virology (1985), 66(4),

719-26

CODEN: JGVIAY; ISSN: 0022-1317

DOCUMENT TYPE: Journal LANGUAGE: English

SOURCE:

DNA complementary to RNA sequences that encode the surface projection (spike) of the coronavirus infectious bronchitis virus (IBV), strain Beaudette, were cloned into pBR322 by using cDNA primed with a specific oligonucleotide. A 5.3 kilobase viral insert in the clone pMB179 was identified. The region of this clone containing the spike gene was sequenced by the chain termination method, and the gene sequence is presented. The sequence of the primary translation product, as deduced from the DNA sequence, is a polypeptide of 1162 amino acids with a mol. weight of 127,006. The polypeptide is subsequently cleaved to S1 and S2, and partial amino acid anal. of the N-terminus of the S1 polypeptide was used to locate the position of this terminus of S1 within the large open reading frame. The amino acid anal. also shows an 18-amino acid putative signal sequence on the primary translation product which is not present on the mature S1 polypeptide.

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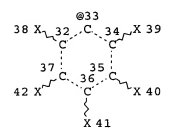
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=> d stat que

L1

STR



VAR G1=C/N/O
VAR G2=O/S/N
VAR G3=O/S
VAR G4=13/19/26/33
NODE ATTRIBUTES:
DEFAULT MLEVEL IS ATOM
DEFAULT ECLEVEL IS LIMITED

GRAPH ATTRIBUTES:

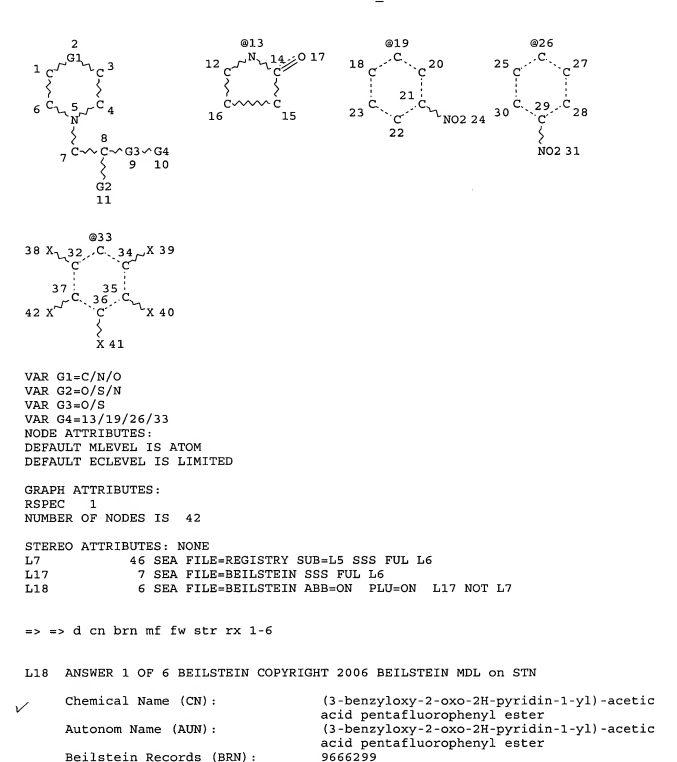
RING(S) ARE ISOLATED OR EMBEDDED

NUMBER OF NODES IS 42

STEREO ATTRIBUTES: NONE

L5 82 SEA FILE=REGISTRY SSS FUL L1

L6 STR



C20 H12 F5 N O4

425.31

Molecular Formula (MF):

Molecular Weight (MW):

```
Reaction:
```

RX

Reaction ID (.ID): 9520825

Reactant BRN (.RBRN): 3617212, 2003848

Reactant (.RCT): (3-benzyloxy-2-oxo-2H-pyridin-1-yl)-acetic

acid, pentafluorophenyl trifluoroacetate

Product BRN (.PBRN): 9666299

Product (.PRO): (3-benzyloxy-2-oxo-2H-pyridin-1-yl)-acetic

acid pentafluorophenyl ester

No. of React. Details (.NVAR): 1

Reaction Details:

RX

Reaction RID (.RID): 9520825.1
Reaction Classification (.CL): Preparation

Yield (.YDT): 3.28 g (BRN=9666299)

Reagent (.RGT): pyridine

Solvent (.SOL): dimethylformamide

Time (.TIM): 1 hour(s)
Temperature (.T): 20 Cel

Reference(s):

 Formica, Mauro; Fusi, Vieri; Giorgi, Luca; Guerri, Annalisa; Lucarini, Simone; Micheloni, Mauro; Paoli, Paola; Pontellini, Roberto; Rossi, Patrizia; Tarzia, Giorgio; Zappia, Giovanni, New J. Chem., CODEN: NJCHE5, 27(11), <2003>, 1575 - 1583; BABS-6432739

Reaction:

RX

Reaction ID (.ID): 9526447

Reactant BRN (.RBRN): 9666299, 3588279

Reactant (.RCT): (3-benzyloxy-2-oxo-2H-pyridin-1-yl)-acetic

acid pentafluorophenyl ester,

1,7-dimethyl-1,4,7,10-tetraazacyclododecane

Product BRN (.PBRN): 9680624

Product (.PRO): 4-(N), 10-(N)-bis<2-(3-benzyloxy-2-oxo-2H-

pyridin-1-yl)acetamido>-1,7-dimethyl-

1,4,7,10-tetraazacyclododecane

No. of React. Details (.NVAR): 1

Reaction Details:

RX

Reaction RID (.RID): 9526447.1
Reaction Classification (.CL): Preparation

Yield (.YDT): 87 percent (BRN=9680624)

Reagent (.RGT): i-Pr2EtN

Solvent (.SOL): dimethylformamide

Time (.TIM): 12 hour(s)
Temperature (.T): 20 Cel

Reference(s):

 Formica, Mauro; Fusi, Vieri; Giorgi, Luca; Guerri, Annalisa; Lucarini, Simone; Micheloni, Mauro; Paoli, Paola; Pontellini, Roberto; Rossi, Patrizia; Tarzia, Giorgio; Zappia, Giovanni, New J. Chem., CODEN: NJCHE5, 27(11), <2003>, 1575 - 1583; BABS-6432739

L18 ANSWER 2 OF 6 BEILSTEIN COPYRIGHT 2006 BEILSTEIN MDL on STN

Chemical Name (CN): 3-methyl-2-<2-oxo-4-(5-oxo-pyrrolidine-2-

carbonyl) -3 - (1-trityl-1H-imidazol-4ylmethyl) -piperazin-1-yl>-butyric acid

pentafluorophenyl ester

Autonom Name (AUN): 3-methyl-2-<2-oxo-4-(5-oxo-pyrrolidine-2-

carbonyl)-3-(1-trityl-1H-imidazol-4ylmethyl)-piperazin-1-yl>-butyric acid

pentafluorophenyl ester

Beilstein Records (BRN): 8605792

Molecular Formula (MF): C43 H38 F5 N5 O5

Molecular Weight (MW): 799.80

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Reaction:
RX
                                     8555989
     Reaction ID (.ID):
     Reactant BRN (.RBRN):
                                      8601073, 1912584
                                      3-methyl-2-<2-oxo-4-(5-oxo-pyrrolidine-2-
     Reactant (.RCT):
                                      carbonyl) -3-(1-trityl-1H-imidazol-4-
                                      ylmethyl)-piperazin-1-yl>-butyric acid,
                                      pentafluorophenol
     Product BRN (.PBRN):
                                      8605792
                                      3-methyl-2-<2-oxo-4-(5-oxo-pyrrolidine-2-
     Product (.PRO):
                                      carbonyl) -3-(1-trityl-1H-imidazol-4-
                                      ylmethyl)-piperazin-1-yl>-butyric acid
                                      pentafluorophenyl ester
     No. of React. Details (.NVAR):
Reaction Details:
     Reaction RID (.RID):
                                      8555989.1
     Reaction Classification (.CL): Preparation
                                     DCC, DMAP, Et3N
     Reagent (.RGT):
     Solvent (.SOL):
                                      CH2Cl2
     Temperature (.T):
                                      0 - 20 Cel
     Reaction Type (.TYP):
                                     Esterification
     Reference(s):
     1. Bhatt, Ulhas; Just, George, Helv.Chim.Acta, CODEN: HCACAV, 83(4),
        <2000>, 722 - 727; BABS-6234236
Reaction:
ВX
                                      8602240
     Reaction ID (.ID):
     Reactant BRN (.RBRN):
                                      8605792
     Reactant (.RCT):
                                      3-methyl-2-<2-oxo-4-(5-oxo-pyrrolidine-2-
                                      carbonyl) -3-(1-trityl-1H-imidazol-4-
                                      ylmethyl)-piperazin-1-yl>-butyric acid
                                      pentafluorophenyl ester
     Product BRN (.PBRN):
                                      8600633
     Product (.PRO):
                                      3-methyl-2-(2-oxo-4-<(5-oxopyrrolidin-2-
                                      yl)carbonyl>-3-<<1-(triphenylmethyl)-1H-
                                      imidazol-4-yl>methyl>piperazin-1-
                                      yl)butanamide
     No. of React. Details (.NVAR):
Reaction Details:
РX
     Reaction RID (.RID):
                                      8602240.1
                                      Preparation
     Reaction Classification (.CL):
     Yield (.YDT):
                                      6.2 mg (BRN=8600633)
     Reagent (.RGT):
     Solvent (.SOL):
                                      ethanol
     Time (.TIM):
                                      8 hour(s)
                                      20 Cel
     Temperature (.T):
                                      Substitution
     Reaction Type (.TYP):
     Reference(s):
     1. Bhatt, Ulhas; Just, George, Helv.Chim.Acta, CODEN: HCACAV, 83(4),
        <2000>, 722 - 727; BABS-6234236
```

(3-methoxymethoxy-2-oxo-2H-pyridin-1-yl)-Autonom Name (AUN): acetic acid 2,5-dioxo-pyrrolidin-1-yl ester 8217191 Beilstein Records (BRN): C13 H14 N2 O7 Molecular Formula (MF): Molecular Weight (MW): 310.26 Reaction: Reaction ID (.ID): 5086550 Reactant BRN (.RBRN): 8202407, 113913 Reactant (.RCT): (3-methoxymethoxy-2-oxo-2H-pyridin-1-yl)acetic acid, N-hydroxy-succinimide Product BRN (.PBRN): 8217191 Product (.PRO): (3-methoxymethoxy-2-oxo-2H-pyridin-1-yl)acetic acid 2,5-dioxo-pyrrolidin-1-yl ester No. of React. Details (.NVAR): Reaction Details: Reaction RID (.RID): 5086550.1 Reaction Classification (.CL): Preparation 22 percent (BRN=8217191) Yield (.YDT): Reagent (.RGT): dicyclohexylcarbodiimide Reference(s): 1. Rai, Bijaya L.; Khodr, Hicham; Hider, Robert C., Tetrahedron, CODEN: TETRAB, 55(4), <1999>, 1129 - 1142; BABS-6157581 L18 ANSWER 4 OF 6 BEILSTEIN COPYRIGHT 2006 BEILSTEIN MDL on STN (3-hydroxy-4-methyl-2-oxo-2H-pyridin-1-yl)-Chemical Name (CN): acetic acid 1,3-dioxo-1,3-dihydro-isoindol-2-yl ester (3-hydroxy-4-methyl-2-oxo-2H-pyridin-1-yl)-Autonom Name (AUN):

RX

RX

2-yl ester

acetic acid 1,3-dioxo-1,3-dihydro-isoindol-

Beilstein Records (BRN): 7944429
Molecular Formula (MF): C16 H12 N2 O6

Molecular Weight (MW): 328.28

```
Reaction:
```

RX

Reaction ID (.ID): 4862410

Reactant BRN (.RBRN): 7920668, 131208

Reactant (.RCT): (3-hydroxy-4-methyl-2-oxo-2H-pyridin-1-yl)-

acetic acid, N-hydroxy-phthalimide

Product BRN (.PBRN): 7944429

Product (.PRO): (3-hydroxy-4-methyl-2-oxo-2H-pyridin-1-yl)-

acetic acid 1,3-dioxo-1,3-dihydro-isoindol-

2-yl ester

No. of React. Details (.NVAR): 1

Reaction Details:

RX

Reaction RID (.RID): 4862410.1
Reaction Classification (.CL): Preparation

Reagent (.RGT): dicyclohexylcarbodiimide

Solvent (.SOL): tetrahydrofuran

Other Conditions (.COND): 0 deg C, 20 min; room temperature, 1 h

Reference(s):

Fox, Raymond C.; Taylor, Paul D., Synth.Commun., CODEN: SYNCAV, 28(9),
 <1998>, 1563-1574; BABS-6089572

Reaction:

RX

Reaction ID (.ID): 4866678

Reactant BRN (.RBRN): 7944429, 1739626

Reactant (.RCT): (3-hydroxy-4-methyl-2-oxo-2H-pyridin-1-yl)-

acetic acid 1,3-dioxo-1,3-dihydro-isoindol-

2-yl ester, tris-(2-amino-ethyl)-amine

Product BRN (.PBRN): 7964315

Product (.PRO): N-<2-(bis-<2-<2-(3-hydroxy-4-methyl-2-oxo-

2H-pyridin-1-yl)-acetylamino>-ethyl>-

amino) -ethyl>-2-(3-hydroxy-4-methyl-2-oxo-

2H-pyridin-1-yl)-acetamide

No. of React. Details (.NVAR): 1

Reaction Details:

RX

Reaction RID (.RID): 4866678.1 Reaction Classification (.CL): Preparation

40 percent (BRN=7964315) Yield (.YDT):

Et3N Reagent (.RGT):

Solvent (.SOL): tetrahydrofuran

Time (.TIM): 1 hour(s)

Ambient temperature Other Conditions (.COND):

Reference(s):

1. Fox, Raymond C.; Taylor, Paul D., Synth.Commun., CODEN: SYNCAV, 28(9), <1998>, 1563-1574; BABS-6089572

L18 ANSWER 5 OF 6 BEILSTEIN COPYRIGHT 2006 BEILSTEIN MDL on STN

(3-hydroxy-2-oxo-2H-pyridin-1-yl)-acetic Chemical Name (CN):

acid 1,3-dioxo-1,3-dihydro-isoindol-2-yl

ester

(3-hydroxy-2-oxo-2H-pyridin-1-yl)-acetic Autonom Name (AUN):

acid 1,3-dioxo-1,3-dihydro-isoindol-2-yl

ester

7944361 Beilstein Records (BRN):

Molecular Formula (MF): C15 H10 N2 O6

Molecular Weight (MW): 314.25

Reaction:

RX

Reaction ID (.ID): 4862403

Reactant BRN (.RBRN): 1453406, 131208

(3-hydroxy-2-oxo-2H-pyridin-1-yl)-acetic Reactant (.RCT):

acid, N-hydroxy-phthalimide

Product BRN (.PBRN): 7944361

Product (.PRO): (3-hydroxy-2-oxo-2H-pyridin-1-yl)-acetic acid 1,3-dioxo-1,3-dihydro-isoindol-2-yl

ester

No. of React. Details (.NVAR):

Reaction Details:

RX

Reaction RID (.RID): 4862403.1 Reaction Classification (.CL): Preparation

Reagent (.RGT): dicyclohexylcarbodiimide

Solvent (.SOL): tetrahydrofuran

Other Conditions (.COND): 0 deg C, 20 min; room temperature, 1 h

Reference(s):

Fox, Raymond C.; Taylor, Paul D., Synth.Commun., CODEN: SYNCAV, 28(9), <1998>, 1563-1574; BABS-6089572

Reaction:

RX

Reaction ID (.ID): 4866677

Reactant BRN (.RBRN): 7944361, 1739626

Reactant (.RCT): (3-hydroxy-2-oxo-2H-pyridin-1-yl)-acetic

acid 1,3-dioxo-1,3-dihydro-isoindol-2-yl

ester, tris-(2-amino-ethyl)-amine

Product BRN (.PBRN): 3643079

Product (.PRO): N-<2-(bis-<2-<2-(3-hydroxy-2-oxo-2H-

pyridin-1-yl)-acetylamino>-ethyl>-amino)ethyl>-2-(3-hydroxy-2-oxo-2H-pyridin-1-yl)-

acetamide

No. of React. Details (.NVAR): 1

Reaction Details:

RX

Reaction RID (.RID): 4866677.1
Reaction Classification (.CL): Preparation

Yield (.YDT): 33 percent (BRN=3643079)

Reagent (.RGT): Et3N

Solvent (.SOL): tetrahydrofuran

Time (.TIM): 1 hour(s)

Other Conditions (.COND): Ambient temperature

Reference(s):

Fox, Raymond C.; Taylor, Paul D., Synth.Commun., CODEN: SYNCAV, 28(9),
 <1998>, 1563-1574; BABS-6089572

L18 ANSWER 6 OF 6 BEILSTEIN COPYRIGHT 2006 BEILSTEIN MDL on STN

Chemical Name (CN): succinimide ester of 1-methyl-1'-

carboxymethyl-4,4'-bipyridinium

perchlorate

Beilstein Records (BRN): 5846197

Molecular Formula (MF): C17 H17 N3 O4 . 2 Cl O4

Molecular Weight (MW): 327.34, 99.45

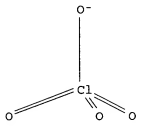
CM 1

FBRN 5832906

FMF C17 H17 N3 O4

CM 2

FBRN 3587878 FMF Cl O4



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Reaction:
RX
     Reaction ID (.ID):
                                      1645997
     Reactant BRN (.RBRN):
                                      5844862, 113913
                                      1-methyl-1'-carboxymethyl-4,4'-
     Reactant (.RCT):
                                      bipyridinium perchlorate,
                                      N-hydroxy-succinimide
                                      5846197
     Product BRN (.PBRN):
     Product (.PRO):
                                      succinimide ester of 1-methyl-1'-
                                      carboxymethyl-4,4'-bipyridinium
                                      perchlorate
     No. of React. Details (.NVAR):
Reaction Details:
RX
     Reaction RID (.RID):
                                      1645997.1
     Reaction Classification (.CL):
                                      Preparation
     Reagent (.RGT):
                                      1,3-dicyclohexylcarbodiimide (DCC)
     Solvent (.SOL):
                                      acetonitrile
     Time (.TIM):
                                      16 hour(s)
     Temperature (.T):
                                      -5 - 20 Cel
     Reference(s):
     1. Tsukahara, Keiichi; Todorobaru, Hiromi, Chem.Lett., CODEN: CMLTAG(7),
        <1992>, 1181-1184; BABS-5704527
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